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Natural Transmission of *Helicobacter saguini* Causes Multigenerational Inflammatory Bowel Disease in C57/129 IL-10−/− Mice

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ABSTRACT Cotton-top tamarins (CTTs) are an ideal model of human inflammatory bowel disease (IBD) because these animals develop multigenerational, lower bowel cancer. We previously isolated and characterized a novel enterohepatic *Helicobacter* species, *Helicobacter saguini*, from CTTs with IBD and documented that *H. saguini* infection in germfree C57BL IL-10−/− mice recapitulates IBD, suggesting that *H. saguini* influences IBD etiopathogenesis. In this study, we utilized a germfree IL-10−/− model to illustrate that *H. saguini* infection can naturally transmit and infect four generations and cause significant intestinal inflammatory pathology. Additionally, whole-genome sequencing of representative *H. saguini* isolates from each generation of IL-10−/− mice revealed gene mutations suggestive of multigenerational evolution. Overall, these results support that specific bacterial species with pathogenic potential, like *H. saguini*, are transmissible microorganisms in the etiopathogenesis of IBD in CTTs and reinforces the importance of specific microbiota in the pathogenesis of IBD in humans.

IMPORTANCE While family history is a significant risk factor for developing inflammatory bowel disease (IBD), it is unclear whether the microbiome from parents is a transmissible influence on disease in their offspring. Furthermore, it is unknown whether IBD-associated microbes undergo genomic adaptations during multigenerational transmission and chronic colonization in their hosts. Herein, we show that a single bacterial species, *Helicobacter saguini*, isolated from a nonhuman primate species with familial IBD, is transmissible from parent to offspring in germfree IL-10−/− mice and causes multigenerational IBD. Additionally, whole-genome sequence analysis of *H. saguini* isolated from different mouse generations identified microevolutions in environmental interaction, nutrient metabolism, and virulence factor genes that suggest that multigenerational transmission may promote adaptations related to colonization and survival in new hosts and chronic inflammatory environments. The findings from our study highlight the importance of specific bacterial species with pathogenic potential, like *H. saguini*, as transmissible microorganisms in the etiopathogenesis of IBD.

KEYWORDS DNA damage, *Helicobacter*, cotton-top tamarins, germfree IL-10−/− mice, inflammatory bowel disease, microbiome, microevolutions, multigenerational, whole-genome sequencing

While remaining idiopathic, inflammatory bowel disease (IBD) is the multifactorial result of genetic, environmental, and microbial interactions. Family history is the greatest risk factor (1, 2). The concordance rate of IBD in monozygotic twins is <50%, indicating incomplete genetic penetrance and reinforcing the importance of shared environmental factors, including the microbiome (3–6). Cotton-top tamarins (*Saguinus oedipus*) (CTTs) are nonhuman primates that, when in captivity, develop multigenera-
tional, idiopathic IBD that progresses to colon cancer (7–10). These animals are considered an ideal model for IBD due to their clinical and histopathological similarities with human IBD; however, they are no longer used because they have been placed on the endangered species list. Interestingly, CTTs are more likely to develop colitis when cohoused in a colony with endemic IBD than when reared apart, suggesting transmission of specific microorganisms, which could be etiologically important (7). Our lab isolated *Helicobacter saguini*, a novel enterohepatic *Helicobacter* species (EHS), from the feces and colons of captive CTTs with IBD (9). Given that EHS infection has been associated with IBD in human patients and macaque species (11, 12) and that other EHS experimentally induce IBD in murine models (13), we hypothesized that *H. saguini* infection is associated with IBD in CTTs. Recently, we demonstrated that monooassociated *H. saguini* infection in germfree IL-10−/− mice caused typhlocolitis and argued that *H. saguini* contributes to IBD in CTTs (14). *H. saguini*, however, was unable to colonize specific-pathogen-free (SPF) IL-10−/− mice (14). In this study, we used the germfree IL-10−/− mouse model to test the hypothesis that *H. saguini* infection in captive CTT colonies is naturally transmitted through successive generations resulting in colitis in infected animals. Additionally, we studied how multigenerational *H. saguini* colonization yielded microevolutions in its genome that may have promoted chronic colonization and the sustained ability of *H. saguini* to induce pathogenicity.

**Experimental results.** To test whether *H. saguini* infection can be naturally transmitted to multiple generations, a male and female breeding pair (F0 generation) were orally dosed and colonized with *H. saguini*. Breeding of F0 yielded three pups (F1), one of which naturally acquired infection after being cohoused with its dam (Fig. 1A). Breeding of the infected F1 female with the F0 male yielded F2, and subsequent brother-sister mating produced F3 followed by F4 that also naturally acquired *H. saguini* infection (Fig. 1A). For all generations, pups remained cohoused with their dams for 5 to 6 weeks after weaning to promote natural transmission of *H. saguini* infection. Infected mice remained *H. saguini* PCR positive at necropsy. Fluorescence *in situ* hybridization (FISH) of the ceca of representative mice from each generation demonstrated that *H. saguini* localized at the mucosal surface, while no bacteria were present in the age-matched control mice (Fig. 1B). These data indicate that *H. saguini* infection chronically colonized germfree IL-10−/− mice for up to ~40 weeks and that infection with *H. saguini* can be naturally transmitted from dams to offspring over four successive generations, likely via fecal-oral horizontal transmission.

Mice PCR positive for *H. saguini* infection in all four generations developed significant inflammatory pathology in the cecum and colon compared to their respective age-matched controls (Fig. 1C; see also Fig. S1A in the supplemental material), except generation F1 in which only 1/3 mice were positive for infection. Litter 3 from generation F4, which failed to naturally acquire infection, did not have statistically different histology scores compared to those of the controls. Pathology in the cecum and colon was characterized by marked infiltration of inflammatory cells in the mucosa and submucosa along with dysplasia, hyperplasia, and epithelial and crypt defects (Fig. 1D and Fig. S1B). Interestingly, male mice from the F3 and F4 generations had considerably higher median cecum and colon pathology scores compared to their female counterparts, although the difference did not reach statistical significance (Fig. 1C and Fig. S1A). Fecal colonization of *H. saguini* was statistically higher in females than in males in the F3 generation and when all generations were pooled (Fig. 1E).

Furthermore, we observed that mice colonized with ≥~10^5 copies of *H. saguini* have cecum and colon pathology scores of <10 and were primarily females, while the opposite trend was seen for males (Fig. 1F and Fig. S1C). This suggests that females may tolerate higher colonization densities and elicit less robust pathology responses to *H. saguini* compared to males in this model.

The levels of inflammatory cytokines for interleukin 22 (IL-22), gamma interferon (IFN-γ), inducible nitric oxide synthase 2 (iNOS2), IL-17a, IL-6, tumor necrosis factor alpha (TNF-α), IL-23a, and IL-4 expressed in the ceca of a representative F4 litter were...
also significantly higher than those for age-matched controls (Fig. 1G). Immunohistochemistry was performed for the DNA histone modification, γ-H2AX, a marker for DNA damage (15) and apoptotic cells (16). *H. saguini*-infected mice from a representative F4 litter demonstrated significantly higher levels of γ-H2AX in the cecum compared to age-matched controls (Fig. S1D and E). These results indicated that multigenerational, chronic *H. saguini* infection induces inflammatory gene pathways as well as DNA damage and cell death.
Genomic microevolutions have been reported for the gastric carcinogen *Helicobacter pylori* during chronic host colonization (17, 18) and transmission to family members (19–21) as well as for the gastroenteritis-causing pathogen *Campylobacter jejuni* (a close relative to EHS) during adaption to new host species (22–26). However, whether microbes associated with IBD undergo genomic microevolution within their hosts is unknown. We sought to address this question by performing comparative genomic analyses between multigenerational *H. saguini* isolates. Representative *H. saguini* isolates were cultured from F0, F2, F3, and F4 mice. Although the mouse from the F1 generation had high *H. saguini* colonization levels (Fig. 1E), an isolate could not be recovered possibly because the organism did not remain viable/culturable after sample collection and frozen storage. Genomes for F0 to F4 isolates were sequenced using Illumina MiSeq and PacBio for high-quality de novo contig assemblies (see Tables S1 and S2 in the supplemental material). Gross rearrangements in chromosomal regions were not detected in the isolates. Variant analysis identified synonymous, nonsynonymous, insertion, and deletion mutational differences between the multigenerational genomes (Table S3) that occurred throughout the chromosome (Fig. 2A). Overall, the number of variants increased in a generational manner (Fig. 2B and C).

![Graphical circular maps of chromosomes from *H. saguini* F0, F2, F3, and F4 isolates.](http://msphere.asm.org)
multigenerational microevolutions that may facilitate colonization and survival in its host. We hypothesized that multigenerational passage of H. saguini in germfree IL-10 mice yields gene variants that would enhance its colonization in SPF IL-10 mice. Therefore, SPF IL-10 mice were inoculated with the H. saguini F4 isolate. However, this strain failed to colonize (data not shown), possibly because the murine microbiota precluded H. saguini colonization, which was similar to our previous study attempting to infect SPF IL-10 mice with this organism (14).

**Conclusion/summary.** Here, we show that H. saguini infection can naturally transmit and persistently colonize several successive generations of germfree IL-10 mice to cause IBD. Given the coprophagous behavior of mice, horizontal transmission via the fecal-oral route is the likely mechanism for transmitting H. saguini infection from the dams to offspring. This is the primary mechanism by which EHS infection, such as H. hepaticus, are also transmitted to new hosts. Whole-genome comparative analyses identified host- and generation-dependent variant genes from H. saguini isolates, suggesting that IBD-associated microbes may adapt for colonization and survival in chronic inflammatory environments. However, in this study, in vitro culturing may have accounted for some genetic changes, although we attempted to minimize this potential by minimally passaging (≤4 times) H. saguini isolates before whole-genome sequencing. Future studies will be required to address the functional significance of the variant genes identified in the multigenerational H. saguini isolates. Nevertheless, our results support the hypothesis that transmission of specific microbiota (i.e., H. saguini) may be important in initiating and maintaining IBD in captive CTTs. In conclusion, the findings of this study warrant future investigations into the role of the microbiome in the etiopathogenesis of multigenerational IBD in other animal models and humans.

**Methods.** Germfree C57BL/6 (B6.129P2-IL-10tm1Cgn [IL-10−/−]) mice were infected with H. saguini MIT 97-6194-4 as previously described (14). Briefly, a male and female breeding pair (F0 generation) received 0.2 ml of fresh inoculum of H. saguini by gastric gavage ever other day for three doses and then were bred to produce the F1 generation. The infected F1 female was bred with the F0 male to yield F2, and brother-sister mating produced the sequential F3 and F4 generations. Pups were cohoused with the dams for 5 to 6 weeks after weaning. Three cohorts of mice were included as age-matched controls. Colonization with H. saguini was confirmed by PCR of feces using the Helicobacter genus-specific primer pair C97/C05. Histopathological evaluation, fluorescence in situ hybridization (FISH), immunohistochemistry for γ-H2AX, and quantitative PCR (qPCR) for cytokine gene expression were performed as previously described (14). Colonization levels of H. saguini in fecal samples were quantified with qPCR targeting the 16s rRNA gene. Genomic copies were calculated from a standard curve of H. saguini genomic DNA in which 1 g of H. saguini genomic DNA (≈2.7 Mb) equals ≈3.43 × 10^{14} molecular copies (assuming 1 bp equals ≈650 g/mol and Avogadro’s number is ≈6.02 × 10^{23} molecules/mol). The levels of eukaryotic 18S rRNA from mouse DNA were determined using qPCR. Colonization levels were then
normalized for each sample by reporting genomic copies of *H. saguini* per µg of mouse 18S rRNA. Cecal and colonic lesion scores, cytokine mRNA expression, *H. saguini* colonization levels, and γ-H2AX expression were analyzed using the Mann-Whitney U nonparametric test. Statistical analyses were performed using GraphPad Prism, version 5.0 (GraphPad Software, Inc., La Jolla, CA). Results were considered significant at a *P* value of <0.05.

*H. saguini* was cultured from fecal samples and intestinal samples as described previously (14). Isolates were repassaged ≤4 times to minimize potential genetic or phenotypic changes. Purified genomic DNA was sequenced by Illumina MiSeq (34) and Pacific Biosciences RSII (35). Spades version 3.10.1 was used for de novo hybrid assembly with the MiSeq reads and the PacBio filtered subreads as long-read scaffolds (36). Resulting contigs were annotated by RAST hosted by PATRIC (37). Sequence variants were identified using the Variation Analysis Service hosted by PATRIC with the BWA-mem aligner and FreeBayes single nucleotide polymorphism (SNP) caller. Further gene annotations and analyses for metabolic function and virulence factors were performed as previously described (27).

**Ethics approval for animal studies.** The protocols for use of animals were approved by the Committee on Animal Care of the Massachusetts Institute of Technology.

**Data availability.** All data and materials generated or analyzed during this study are included in this published article and its supplemental material files or are available from the corresponding author(s) upon request. DNA sequence and assembled genome data have been deposited in NCBI under the BioProject accession no. PRJNA449717.

**SUPPLEMENTAL MATERIAL**

Supplemental material is available online only.

**FIG S1**, TIF file, 2.6 MB.

**TABLE S1**, DOCX file, 0.01 MB.

**TABLE S2**, DOCX file, 0.01 MB.

**TABLE S3**, XLSX file, 0.2 MB.

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We declare that we have no competing interests.

A.M. performed animal necropsies, tissue processing, bacterial culture, DNA isolation, genome assembly and analysis, data analysis and statistics, and wrote the manuscript. Z.S. performed tissue processing and bacterial culture, interpreted data, and reviewed the manuscript. Y.F. performed tissue processing for immunohistochemistry (IHC). D.G. performed animal necropsies and tissue processing. S.M. performed histopathological analysis. M.T.W. supervised germfree facility studies. J.G.F. conceived the project, supervised all studies, interpreted data, and reviewed and edited the manuscript. All authors read and approved the final manuscript.

**REFERENCES**

1. Russell RK, Satsangi J. 2008. Does IBD run in families? Inflamm Bowel Dis 14(Suppl 2):S20 –S21. https://doi.org/10.1002/ibd.20573.

2. Halme L, Paavola-Sakki P, Turunen U, Lappalainen M, Farkkila M, Kontula K. 2006. Family and twin studies in inflammatory bowel disease. World J Gastroenterol 12:3668 –3672. https://doi.org/10.3748/wjg.v12.i23.3668.

3. Albenberg LG, Lewis JD, Wu GD. 2012. Food and the gut microbiota in inflammatory bowel diseases: a critical connection. Curr Opin Gastroenterol 28:314 –320. https://doi.org/10.1097/MOG.0b013e328354586f.

4. Lane ER, Zisman TL, Suskind DL. 2017. The microbiota in inflammatory bowel disease: current and therapeutic insights. J Inflamm Res 10:63–73. https://doi.org/10.2147/JIR.S116088.

5. Ni J, Wu GD, Albenberg L, Tomov VT. 2017. Gut microbiota and IBD: causation or correlation? Nat Rev Gastroenterol Hepatol 14:573–584. https://doi.org/10.1038/nrgastro.2017.88.

6. Lepage P, Hasler R, Spehlmann ME, Rehman A, Zvirbliene A, Begun A, Ott S, Kupcinskas L, Dore J, Raedler A, Schreiber S. 2011. Twin study indicates loss of interaction between microbiota and mucosa of patients with ulcerative colitis. Gastroenterology 141:227–236. https://doi.org/10.1053/j.gastro.2011.04.011.

7. Johnson LD, Ausman LM, Sehgal PK, King N. 1996. A prospective study of the epidemiology of colitis and colon cancer in cotton-top tamarins (Saguinus oedipus). Gastroenterology 110:102–115. https://doi.org/10.1053/gast.1996.v110.pm8536845.

8. Wood JD, Peck OC, Tefend KS, Rodriguez MM, Rodriguez MV, Hernandez PR, et al. 2017. Gut microbiota and IBD: association or correlation? Nat Rev Gastroenterol Hepatol 14:573–584. https://doi.org/10.1038/nrgastro.2017.88.
dez CJ, Stonerook MJ, Sharma HM. 1998. Colitis and colon cancer in cotton-top tamarins (Saguinus oedipus oedipus) living wild in their natural habitat. Dig Dis Sci 43:1443–1453. https://doi.org/10.1023/a:1018842103300.

9. Saunders KE, Shen Z, Dewhirst FE, Paster BJ, Dangler CA, Fox JG. 1999. Novel intestinal Helicobacter species isolated from cotton-top tamarins (Saguinus oedipus) with chronic colitis. J Clin Microbiol 37:146–151. https://doi.org/10.1128/JCM.37.1.146-151.1999.

10. Bertone ER, Giovannucci EL, King NWI, Petto AJ, Johnson LD. 1998. Family history as a risk factor for ulcerative colitis-associated colon cancer in cotton-top tamarin. Gastroenterology 114:669–674. https://doi.org/10.1016/S0016-5085(98)70580-3.

11. Castano-Rodriguez N, Kaakouch NO, Lee WS, Mitchell HM. 2017. Dual role of Helicobacter and Campylobacter species in IBD: a systematic review and meta-analysis. Gut 66:235–249. https://doi.org/10.1136/gutjnl-2015-310545.

12. Hansen R, Thomson JM, Fox JG, El-Omar EM, Hold GL. 2011. Could Helicobacter organisms cause inflammatory bowel disease? FEMS Immunol Med Microbiol 61:1–14. https://doi.org/10.1111/j.1574-695X.2010.00744.x.

13. Whary MT, Fox JG. 2004. Natural and experimental Helicobacter infections. Comp Med 54:128–158.

14. Shen Z, Mannion A, Whary MT, Muthupalani S, Sheh A, Feng Y, Gong G, Vandamme P, Holcombe HR, Paster BJ, Fox JG. 2016. Helicobacter saguini, a novel helicobacter isolated from cotton-top tamarins with ulcerative colitis, has proinflammatory properties and induces typhlocolitis and dysplasia in gnotobiotic IL-10−/− mice. Infect Immun 84: 2307–2316. https://doi.org/10.1128/IAI.00235-16.

15. Rogakou EP, Pilch DR, Orr AH, Ivanova VS, Bonner WM. 1998. DNA double-stranded breaks induce histone H2AX phosphorylation on serine 139. J Biol Chem 273:5858–5868. https://doi.org/10.1074/jbc.273.10.5858.

16. Rogakou EP, Nieves-Neira W, Boon C, Pommier Y, Bonner WM. 2000. Initiation of DNA fragmentation during apoptosis induces phosphorylation of H2AX histone at serine 139. J Biol Chem 275:4818–4822. https://doi.org/10.1074/jbc.275.13.4818.

17. Lundin A, Bjorkholm B, Kupershmidt I, Uemo No, Nilsson P, Andersson DI, Engstrand L. 2005. Slow genetic divergence of Helicobacter pylori strains during long-term colonization. Infect Immun 73:9390–9395. https://doi.org/10.1128/IAI.73.11.9390-9395.2005.

18. Nell S, Kennesmann L, Schwarz S, Josenhaus C, Suerbaum S. 2014. Dynamics of Lewis b binding and sequence variation of the babA protein of Campylobacter jejuni strains during long-term colonization. Infect Immun 82:430–438. https://doi.org/10.1128/IAI.00750-14.

19. Didelot X, Nell S, Yang J, Woltemate S, van der Merwe S, Suerbaum S. 2013. Genomic evolution and transmission of Helicobacter pylori in two South African families. Proc Natl Acad Sci USA 110:13880–13885. https://doi.org/10.1073/pnas.1304681110.

20. Morelli G, Didelot X, Kusecek B, Schwarz S, Bahlawane C, Falush D, Suerbaum S, Achtman M. 2013. Microevolution of Helicobacter pylori during prolonged infection of single hosts and within families. PLoS Genet 6:e1001036. https://doi.org/10.1371/journal.pgen.1001036.

21. Furuta Y, Konno M, Osaki T, Yonezawa H, Ishige T, Imai M, Shiwa Y, Shibata-Hatta M, Kanasaki Y, Yoshikawa H, Kamiya S, Kobayashi I. 2015. Microevolution of virulence-related genes in Helicobacter pylori familial infection. PLoS One 10:e0127197. https://doi.org/10.1371/journal.pone.0127197.

22. Jerome JP, Bell JA, Plovanich-Jones AE, Barrick JE, Brown CT, Mansfield LS. 2011. Standing genetic variation in contingency loci drives the rapid microevolution of virulence-related genes in Helicobacter pylori lipopolysaccharide due to changes in the lengths of poly(C) tracts in alpha3-fucosyltransferase genes. Infect Immun 79:3561–3566. https://doi.org/10.1128/IAI.00235-16.

23. Vandenbroucke-Grauls CM, Kusters JG. 1999. Phase variation in Helicobacter pylori lipopolysaccharide. Microbiology 145:2211–2216. https://doi.org/10.1099/0016-5365-145-9-2211.

24. Mannion A, Fabian N, Stair M, Dzink-Fox J, Carrasco SE, Buckley-Jordan E, Annamalai D, Fox JG. 2019. Draft genome sequences of Klebsiella pneumoniae strains isolated from immunocompromised NOD-scid gamma research mice. Microbiol Resour Announc 8:e00869-19. https://doi.org/10.1128/MRA.00869-19.

25. Mannion AJ, Martin HR, Shen Z, Buckley EM, Dzink-Fox JL, Garcia A, Martini RP, Patterson MM, Fox JG. 2018. Plasmid-mediated quinolone resistance in Shigella flexneri isolated from macaques. Front Microbiol 9:311. https://doi.org/10.3389/fmicb.2018.00311.

26. Nork S, Bankевич A, Antipov D, Gurevich AA, Korobeynikov A, Lapidus A, Pribjalski AD, Pysbin A, Sirotkin A, Sirotkin Y, Stepanauskas R, Clingenpeel SR, Woyke T, Mclean JS, Lasken R, Tesler G, Alekseyev MA, Pevzner PA. 2013. Assembling single-cell genomes and metagenomes from chimeric MDA products. J Comput Biol 20:714–737. https://doi.org/10.1089/cmb.2013.0084.

27. Wattam AR, Davis JJ, Assaf R, Boisvert S, Brettin T, Bun C, Conrad N, Dietrich EM, Diz T, Gabbard JL, Gerdes S, Henry CS, Kenyon RW, Machi D, Mao C, Nordberg EK, Olsen GJ, Murphy-OLson DE, Olson R, Overbeek R, Plovanich-Jones A, Prjibelski AD, Pyshkin A, Sirotkin A, Stepanauskas R, Clin-genpeel SR, Woyke T, McLean JS, Lasken R, Tesler G, Alekseyev MA, Pevzner PA. 2013. Assembling single-cell genomes and metagenomes from chimeric MDA products. J Comput Biol 20:714–737. https://doi.org/10.1089/cmb.2013.0084.

28. Nakamura H, Avalashvili P, Frade S, Setit A, Jensen C, Westlund A, Gotfredsen P, Bard J, Vesterlund S. 2020. Improved improvements. Improvements of PATRC, the all-bacterial Bioinformatics Database and Analysis Resource Center. Nucleic Acids Res 45: D535–D542. https://doi.org/10.1093/nar/gkw1017.