Acid-suppressive medications and risk of colorectal cancer: results from three large prospective cohort studies

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BACKGROUND: Despite several plausible biological mechanisms linking proton pump inhibitors (PPIs) and H2 receptor antagonists (H2RAs) with colorectal tumorigenesis, their association with risk of colorectal cancer (CRC) has not been adequately assessed in prospective epidemiological studies.

METHODS: We evaluated the association of acid-suppressive medication use with CRC risk among 175,871 (PPI) and 208,831 (H2RA) participants from three large prospective cohort studies. Medication use was assessed at baseline and updated biennially. The association was evaluated using multivariate Cox proportional hazards regression models.

RESULTS: There was no significant association between baseline PPI use (hazard ratio (HR) = 0.89, 95% confidence interval (CI), 0.71–1.12) or PPI use after a lag of 8–10 years (HR = 1.12, 95% CI, 0.78–1.59) with CRC risk. We observed no significant association between H2RA use after a lag of 8–10 years and CRC risk (HR = 1.02, 95% CI, 0.81–1.28), while risk was lower for participants with baseline H2RA use (HR = 0.76, 95% CI, 0.60–0.95). Duration of PPI use or H2RA use was not associated with CRC risk (P-trend = 0.21 and 0.95, respectively).

CONCLUSIONS: Among participants from three large prospective cohorts, use of PPI or H2RA was not associated with higher risk of colorectal cancer.

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Published online: 16 June 2020

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Received: 7 January 2020 Revised: 5 May 2020 Accepted: 28 May 2020

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British Journal of Cancer (2020) 123:844–851; https://doi.org/10.1038/s41416-020-0939-y
a prospective analysis to evaluate the association between use of acid-suppressive medications and colorectal cancer risk within three large cohorts (Nurses’ Health Study [NHS], Nurses’ Health Study II [NHSII] and Health Professionals Follow-Up Study [HPFS]) with updated information on PPI and H2RA use, with up to 26 years of follow-up.

METHODS
Study population
The NHS, NHSII and HPFS are ongoing US-based prospective cohort studies. The NHS was established in 1976 among 121,700 US female registered nurses aged 30–55 years,29,30 and NHSII was established in 1989 among 116,429 US female nurses aged 25–42 years.31 HPFS was established in 1986 among 51,529 US male health professionals aged 40–75 years at baseline.32 In all three cohorts, participants completed an initial questionnaire about their medical history, lifestyle factors and health behaviours. Participants have been mailed follow-up questionnaires every 2 years to provide updated information on those factors, as well as disease outcomes. Participants were also asked to complete semi-quantitative validated food frequency questionnaires (FFQs) approximately every 4 years. Follow-up is >90% for all three cohorts. The study protocol was approved by the institutional review boards of the Brigham and Women’s Hospital and Harvard T.H. Chan School of Public Health, and those of participating registries as required.

The eligible population for the analysis of PPI use consisted of participants who answered the questionnaire in which PPI use was first assessed (NHS, 2000; NHSII, 2001; HPFS, 2004). We excluded participants with previous history of cancer, except non-melanoma skin cancer and history of ulcerative colitis or Crohn’s disease. After those exclusions, a total of 175,871 participants were included in the analysis. Among eligible participants who answered the first questionnaire where H2RA use was assessed (NHS, 1994; NHSII, 1995; HPFS, 1998), we applied the same exclusions, leaving a total of 208,831 participants.

Ascertainment of PPI and H2RA use
Information about PPI use and H2RA use was obtained from the baseline questionnaire, and updated in each subsequent biennial cycle. At baseline, participants were asked about regular use (yes/no) of Prilosec or Prevacid in NHS and NHSII, and Prilosec, Nexium, Prevacid (lansoprazole), Protonix or Aciphex in HPFS in the past 2 years. Similarly, they were asked about regular use of ranitidine (Zantac) or cimetidine (Tagamet) in NHS, cimetidine (Tagamet) or other H2 blockers (e.g. Zantac, Pepcid, Axid) in NHSII, and cimetidine or ranitidine (e.g. Tagamet, Zantac) in HPFS in the past 2 years. If a participant’s information on medication use was missing for a follow-up cycle, then that participant did not contribute person-time to the analysis for that cycle.

Ascertainment of covariates
Information about demographic and lifestyle factors known to be associated with colorectal cancer was obtained from the baseline and follow-up questionnaires. These included age, height, weight, physical activity, family history of colorectal cancer, regular use of aspirin and non-aspirin nonsteroidal anti-inflammatory drugs (NSAIDs), menopausal status, menopausal hormone therapy (MHT) use, multivitamin use and history of lower endoscopy (including indication for the procedure). All the covariates were updated biennially, except for family history of colorectal cancer, which was updated approximately every 4 years. Once a participant reported an endoscopy, positive history of endoscopy was carried forward in all the subsequent cycles. Information on dietary factors, including intake of fibre, vitamin D, calcium, folate, red meat, processed meat, total fat intake and total caloric intake, was extracted from FFQs every 4 years. We used energy-adjusted values and cumulative updated intake, defined as the average intake from all available FFQs up to the most recent biennial follow-up questionnaire. Participants were asked about a history of gastric or duodenal ulcer in NHS only at two time points (2000, 2008); in the other two cohorts, this information was updated every 2 years after the baseline (1991 in NHSII, 1986 in HPFS).

Documentation of colorectal cancer cases and deaths
Information about disease outcomes, including colorectal cancer, was obtained in biennial questionnaires. For each participant reporting a colorectal cancer diagnosis, we asked for permission to obtain medical records and pathology reports. Deaths of participants were identified through family members, National Death Index or the US postal service. If the review of death certificate revealed colorectal cancer as the cause of death, we sought permission from next of kin to obtain and review medical records. A study physician blinded to exposure status reviewed medical records, and extracted information on cancer stage, histopathologic features and tumour location.

Statistical analysis
Due to the similar questionnaire design across cohorts, we combined data from all three cohorts into a single dataset. Person-time was calculated as time from the return of baseline questionnaires until colorectal cancer diagnosis, diagnosis of any cancer, except non-melanoma skin cancer, death or the end of follow-up (NHS: June 2014; NHSII: June 2015; HPFS: January 2014), whichever came first.

To consider different intervals between exposures and outcome, we used several approaches to modelling exposures, including current, baseline, persistent, cumulative and lagged use of PPI and H2RA. Current use was defined as regular medication use reported in the most recent questionnaire; this variable was updated in each biennial follow-up cycle. Baseline use was assessed on the first questionnaire that asked about PPI or H2RA use, and subsequently carried forward. Persistent use was defined as use reported in the baseline questionnaire plus one or two subsequent follow-up questionnaire cycles. To evaluate the cumulative effect of drug use on cancer risk, we assessed the number of cycles with reported drug use (no use, use in 1–2 cycles, use in ≥3 cycles), regardless of whether these cycles were consecutive or not. Lastly, since the association between several established risk factors and colorectal cancer is observed only after a latency of 10 years or longer,33,34 we examined medication use lagged for varying periods of time up to 10 years. For example, in NHS, PPI use was first assessed in 2000. For a latency of 2–4 years, we used PPI use in 2000 for the follow-up period from 2002 to 2004, PPI use in 2002 for the follow-up period from 2004 to 2006 and so forth. For a latency of 4–6 years, we used PPI use in 2000 for the follow-up period from 2004 to 2006, PPI use in 2002 for the follow-up period from 2006 to 2008 and so on.

We used Cox proportional hazards regression models stratified by age (months) and 2-year questionnaire cycle to estimate the hazard ratios (HRs) and 95% confidence intervals (CIs) of cancer risk for users of acid-suppressive medications, compared to non-users. In the main multivariate model, we adjusted for study cohort (NHS, NHSII, HPFS) and established colorectal cancer risk factors, including BMI (<25 kg/m2, 25–27.5 kg/m2, 27.5–30 kg/m2, ≥30 kg/m2), physical activity (<3 metabolic equivalent [MET]-h/week, 3–27 MET-h/week, ≥27 MET-h/week), family history of colorectal cancer (no, yes), daily caloric intake (quintiles), alcohol intake (<5 g/day, 5–15 g/day, ≥15 g/day), pack-years of smoking (0, 1–10, >10 pack-years), history of lower endoscopy (never, ever), dietary and supplemental vitamin D intake (quintiles), dietary and supplemental calcium intake (quintiles), regular aspirin use (no, yes), dietary and supplemental folate intake (quintiles), MHT use (premenopausal, postmenopausal never MHT user, postmenopausal past MHT user, postmenopausal current MHT user) (NHS and

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Table 1. Baseline characteristics of participants by use of proton pump inhibitors.

| Characteristic\* | Nurses' Health Study (2000) | Nurses’ Health Study II (2001) | Health Professionals Follow-Up Study (2004) |
|------------------|-----------------------------|-------------------------------|--------------------------------------------|
|                  | Non-users (n = 141,809)     | Users (n = 9368)              | Non-users (n = 20,845)                     | Users (n = 3,837)                      |
| Age (years)b     | 56 (11)                     | 57 (11)                       | 69 (9)                                     | 70 (9)                                 |
| Body mass index (kg/m²) | 25 (5)                     | 27 (6)                        | 26 (3)                                     | 26 (3)                                 |
| Activity (MET-h/week) | 19 (19)                    | 16 (16)                       | 35 (28)                                    | 32 (26)                                |
| Total folate intake (µg/day) | 462 (212)                 | 458 (205)                     | 609 (231)                                  | 623 (233)                              |
| Total vitamin D (IU/day) | 372 (201)                 | 377 (198)                     | 446 (228)                                  | 461 (234)                              |
| Total calcium intake (mg/day) | 1058 (388)                | 1081 (403)                    | 969 (339)                                  | 992 (355)                              |
| Total fat (g/day) | 60 (10)                     | 60 (10)                       | 68 (12)                                    | 69 (12)                                |
| Total fibre (g/day) | 19 (5)                     | 18 (5)                        | 23 (6)                                     | 22 (6)                                 |
| Alcohol (g/day)  | 5 (7)                       | 4 (7)                         | 11 (13)                                    | 11 (13)                                |
| Total energy intake (kcal/day) | 1767 (465)                | 1778 (473)                    | 2000 (531)                                 | 1989 (534)                             |
| Red meat (servings/week) | 4 (2)                     | 4 (2)                         | 4 (3)                                      | 4 (3)                                  |
| Processed meat (servings/week) | 2 (2)                   | 2 (2)                         | 2 (2)                                      | 2 (2)                                  |
| History of sigmoidoscopy/colonoscopy | 39 | 55 | 73 | 84 |
| Family history of colorectal cancer | 13 | 14 | 13 | 14 |
| H2RA use | 6 | 11 | 5 | 14 |
| Regular aspirin use (≥2 tablets/week) | 27 | 35 | 49 | 53 |
| Regular NSAID use (≥2 tablets/week) | 33 | 44 | 29 | 39 |
| Multivitamin use | 62 | 63 | 68 | 71 |
| White | 97 | 98 | 96 | 97 |
| Menopausal hormone therapy use | 40 | 48 | N/A | N/A |
| Postmenopausal status | 55 | 59 | N/A | N/A |
| History of smoking | \* | \* | \* | \* |
| Never | 56 | 52 | 53 | 49 |
| Past | 35 | 39 | 44 | 48 |
| Current | 9 | 9 | 4 | 3 |

H2RA H2 receptor antagonist, MET metabolic equivalent, NSAID nonsteroidal anti-inflammatory drugs.

\*Age-adjusted means (standard deviation) for continuous variables, and percentages for categorical variables.

\*Values not age adjusted.

RESULTS

We documented 1255 colorectal cancer cases during 2.1 million years of follow-up in the PPI use analysis, and 2413 cases during 3.8 million years of follow-up in the H2RA use analysis.

Compared to their respective non-users, both PPI and H2RA users had on average a higher BMI, lower levels of physical activity and higher total calcium intake. Users were more likely to have a history of lower endoscopy, in particular due to gastrointestinal symptoms, and use aspirin, NSAID and multivitamins (Tables 1 and 2). Among women, PPI and H2RA users were more likely to be postmenopausal and MHT users. There were fewer never smokers among both PPI and H2RA users.

In multivariate adjusted models, there was no significant association between baseline use of PPI and risk of colorectal cancer (HR = 0.89, 95% CI, 0.71–1.12), while current PPI use was associated with decreased risk (HR = 0.82, 95% CI, 0.68–0.98) (Table 3). Persistent PPI use defined as use in the baseline, and one (HR = 0.83, 95% CI, 0.58–1.18) or two subsequent cycles (HR = 0.84, 95% CI, 0.56–1.24) was not significantly associated with risk (Table 3). We next evaluated cumulative PPI use, and observed no significant association between number of cycles with reported PPI use and risk of colorectal cancer (P-trend = 0.21) (Table 4). To examine different latency periods between PPI use and colorectal cancer diagnosis, we lagged PPI use for up to 10 years, and observed no significant association (8–10 years lag: HR = 1.12, 95% CI, 0.78–1.59) (Table 5). Use of H2RA at baseline was associated with decreased risk of colorectal cancer (HR = 0.76, 95% CI, 0.60–0.95), while there was no significant association between current or persistent H2RA use (Table 3). Duration of H2RA use was also not significantly associated with risk (P-trend = 0.95) (Table 4). We next examined several latency periods up to 10 years, and observed no significant association (8–10 years lag: HR = 1.02, 95% CI, 0.81–1.28) (Table 5).

The associations between PPI use and H2RA use and colorectal cancer risk each remained materially unchanged after adjusting...
for additional potential confounders, including waist circumference, waist-to-hip ratio, NSAID use, intake of processed meat and history of gastric or duodenal ulcer (data not shown). Associations between both medications and risk of colorectal cancer were similar by primary tumour location (Supplementary Table 1) and by study cohort (P-heterogeneity ≥0.13). Furthermore, the associations for both medications and colorectal cancer risk were similar in analyses stratified by sex, history of lower endoscopy, BMI and smoking status (all P-interaction ≥0.24) (Supplementary Table 2). Since PPI users were also more likely to use H2RA (Table 1), we adjusted simultaneously for use of both medications to examine their independent association with colorectal cancer. Neither baseline use of PPI (HR = 0.95, 95% CI, 0.75–1.21) nor H2RA (HR = 0.82, 95% CI, 0.59–1.12) was associated with risk of colorectal cancer in these models.

**DISCUSSION**

In this pooled analysis of three large prospective US cohorts, we found no evidence of overall significant association between use of PPI or H2RA medications and risk of colorectal cancer.

Use of both PPI and H2RA was examined using several approaches, with varying latency periods between medication use and colorectal cancer diagnosis. In the analysis of PPI use, we observed no significant association with baseline, persistent, cumulative or up to 10-year lagged medication use and colorectal cancer risk. We observed decreased colorectal cancer risk in current users of PPI; however, use in the past 2 years before diagnosis is not biologically likely to play a role in disease development, and is more likely to be affected by subclinical disease. There was no significant change in colorectal cancer risk in current H2RA users. We did observe a 23% decrease in colorectal cancer risk that was statistically significant among baseline H2RA users. However, we observed no association with persistent use or longer duration use, arguing against the inverse association between H2RA and colorectal cancer being causal. Overall, these observations do not support the causal role of PPI or H2RA in colorectal cancer aetiology.

Several case–control studies have examined PPI use in relation to colorectal cancer risk, and reported either no association or moderately increased risk among PPI users. One case–control study reported no association between H2RA use and risk of colorectal cancer. While most of those studies relied on information about medication use extracted from health insurance records or medical charts; therefore, eliminating recall bias, over-the-counter use of these medications was not considered, which could lead to exposure misclassification. Furthermore, several studies failed to account for important confounders such as BMI or history of colonoscopy. In our study, users of PPI and H2RA had a higher BMI and were more likely to have a history of colonoscopy, which have been associated with increased or decreased risk of CRC, respectively. Most importantly, in previous studies timing of medication use in relation to colorectal cancer diagnosis was not carefully examined. Colorectal cancer

| Characteristic* | Nurses’ Health Study (1994) | Nurses’ Health Study II (1995) | Health Professionals Follow-up Study (1988) |
|----------------|-----------------------------|------------------------------|-------------------------------------------|
|                | Non-users (n = 162,473) | Users (n = 9001) | Non-users (n = 36,433) | Users (n = 1008) |
| Age (years)
  | 50 (12) | 52 (11) | 56 (10) | 59 (10) |
| Body mass index (kg/m²) | 25 (5) | 27 (6) | 26 (3) | 26 (3) |
| Activity (MET·h/week) | 20 (24) | 18 (22) | 25 (27) | 20 (22) |
| Total folate intake, (µg/day) | 441 (211) | 438 (210) | 480 (273) | 461 (261) |
| Total vitamin D (IU/day) | 364 (211) | 374 (220) | 405 (307) | 406 (279) |
| Total calcium intake (mg/day) | 993 (375) | 1023 (398) | 902 (418) | 909 (409) |
| Total fat (g/day) | 60 (10) | 61 (10) | 71 (14) | 72 (14) |
| Total fibre (g/day) | 18 (5) | 18 (5) | 21 (7) | 20 (6) |
| Alcohol (g/day) | 4 (7) | 4 (7) | 11 (15) | 12 (15) |
| Total energy intake (kcal/day) | 1760 (473) | 1787 (492) | 1995 (616) | 2040 (621) |
| Red meat (servings/week) | 4 (2) | 4 (3) | 4 (3) | 5 (4) |
| Processed meat (servings/week) | 2 (2) | 2 (2) | 3 (3) | 3 (3) |
| History of sigmoidoscopy/colonoscopy | 25 | 37 | 24 | 32 |
| Family history of colorectal cancer | 10 | 11 | 9 | 9 |
| Regular aspirin use (≥2 tablets/week) | 23 | 31 | 41 | 44 |
| Regular NSAID use (≥2 tablets/week) | 26 | 43 | 10 | 21 |
| Multivitamin use | 44 | 45 | 41 | 44 |
| White | 97 | 97 | 96 | 97 |
| Menopausal hormone therapy use | 29 | 37 | N/A | N/A |
| Postmenopausal status | 43 | 47 | N/A | N/A |
| History of smoking | | | | |
| Never | 56 | 51 | 49 | 40 |
| Past | 32 | 35 | 42 | 49 |
| Current | 12 | 14 | 8 | 12 |

MET: metabolic equivalent, NSAID: nonsteroidal anti-inflammatory drugs.
*Age-adjusted means (standard deviation) for continuous variables, and percentages for categorical variables.
1Values not age adjusted.

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Table 3. Association between use of acid-suppressive medications and risk of colorectal cancer.

| Proton pump inhibitors use | Non-users | Users | Non-users | Users | Non-users | Users |
|----------------------------|-----------|-------|-----------|-------|-----------|-------|
| Baseline use               |           |       |           |       |           |       |
| No. of cases               | 998       | 53    | 174       | 30    | 1172      | 83    |
| Person-years               | 1,785,384 | 115,921 | 172,283  | 31,207 | 1,957,667 | 147,128 |
| Age-adjusted HR (95% CI)   | Ref.      | 0.78 (0.59–1.02) | Ref. | 0.92 (0.62–1.36) | Ref. | 0.84 (0.67–1.04) |
| Multivariate HR (95% CI)   | Ref.      | 0.86 (0.65–1.14) | Ref. | 1.02 (0.68–1.52) | Ref. | 0.89 (0.71–1.12) |
| Use in two consecutive baseline cycles |           |       |           |       |           |       |
| No. of cases               | 1027      | 20    | 187       | 13    | 1214      | 33    |
| Person-years               | 1,831,700 | 53,137 | 182,512  | 17,897 | 2,014,212 | 71,034 |
| Age-adjusted HR (95% CI)   | Ref.      | 0.77 (0.49–1.20) | Ref. | 0.71 (0.40–1.25) | Ref. | 0.76 (0.55–1.10) |
| Multivariate HR (95% CI)   | Ref.      | 0.87 (0.55–1.35) | Ref. | 0.81 (0.45–1.44) | Ref. | 0.83 (0.58–1.18) |
| Use in three consecutive baseline cycles |           |       |           |       |           |       |
| No. of cases               | 1030      | 16    | 190       | 10    | 1220      | 26    |
| Person-years               | 1,839,152 | 43,172 | 186,371  | 13,010 | 2,025,523 | 56,182 |
| Age-adjusted HR (95% CI)   | Ref.      | 0.75 (0.46–1.23) | Ref. | 0.74 (0.39–1.41) | Ref. | 0.77 (0.52–1.14) |
| Multivariate HR (95% CI)   | Ref.      | 0.85 (0.52–1.40) | Ref. | 0.84 (0.44–1.62) | Ref. | 0.84 (0.56–1.24) |
| Current use                |           |       |           |       |           |       |
| No. of cases               | 873       | 111   | 159       | 26    | 1032      | 137   |
| Person-years               | 1,472,450 | 245,294 | 152,585  | 28,702 | 1,625,035 | 273,996 |
| Age-adjusted HR (95% CI)   | Ref.      | 0.72 (0.59–0.88) | Ref. | 0.83 (0.54–1.27) | Ref. | 0.73 (0.61–0.88) |
| Multivariate HR (95% CI)   | Ref.      | 0.80 (0.65–0.98) | Ref. | 0.91 (0.59–1.41) | Ref. | 0.82 (0.68–0.98) |
| H2 receptor antagonists use |           |       |           |       |           |       |
| Baseline use               |           |       |           |       |           |       |
| No. of cases               | 1486      | 56    | 837       | 22    | 2323      | 78    |
| Person-years               | 2,881,874 | 153,014 | 705,765  | 17,711 | 3,587,639 | 170,724 |
| Age-adjusted HR (95% CI)   | Ref.      | 0.66 (0.51–0.87) | Ref. | 0.96 (0.63–1.48) | Ref. | 0.71 (0.56–0.88) |
| Multivariate HR (95% CI)   | Ref.      | 0.70 (0.53–0.92) | Ref. | 1.01 (0.65–1.55) | Ref. | 0.76 (0.60–0.95) |
| Use in two consecutive baseline cycles |           |       |           |       |           |       |
| No. of cases               | 1495      | 42    | 771       | 11    | 2266      | 53    |
| Person-years               | 2,928,762 | 91,273 | 647,808  | 7706   | 3,576,570 | 98,979 |
| Age-adjusted HR (95% CI)   | Ref.      | 0.80 (0.59–1.09) | Ref. | 1.12 (0.61–2.05) | Ref. | 0.81 (0.62–1.07) |
| Multivariate HR (95% CI)   | Ref.      | 0.85 (0.62–1.16) | Ref. | 1.24 (0.67–2.27) | Ref. | 0.90 (0.68–1.18) |
| Use in three consecutive baseline cycles |           |       |           |       |           |       |
| No. of cases               | 1213      | 20    | 699       | 7     | 1912      | 27    |
| Person-years               | 1,285,353 | 29,968 | 607,977  | 5463   | 1,893,330 | 35,430 |
| Age-adjusted HR (95% CI)   | Ref.      | 0.69 (0.44–1.07) | Ref. | 0.96 (0.45–2.05) | Ref. | 0.72 (0.49–1.05) |
| Multivariate HR (95% CI)   | Ref.      | 0.73 (0.47–1.14) | Ref. | 1.08 (0.51–2.31) | Ref. | 0.79 (0.54–1.15) |
| Current use                |           |       |           |       |           |       |
| No. of cases               | 1294      | 102   | 699       | 45    | 1993      | 147   |
| Person-years               | 2,333,665 | 173,958 | 603,823  | 31,002 | 2,937,488 | 204,960 |
| Age-adjusted HR (95% CI)   | Ref.      | 0.94 (0.77–1.15) | Ref. | 1.15 (0.85–1.56) | Ref. | 0.98 (0.83–1.16) |
| Multivariate HR (95% CI)   | Ref.      | 1.00 (0.81–1.22) | Ref. | 1.23 (0.90–1.67) | Ref. | 1.06 (0.89–1.26) |

BMI: body mass index, CI: confidence interval, HR: hazards ratio, MHT: menopausal hormone therapy, NHS: Nurses' Health Study, NHSII: Nurses' Health Study II.

*Adjusted for age, BMI (<25 kg/m², 25–27.5 kg/m², 27.5–30 kg/m², ≥30 kg/m²), physical activity (<3 MET-h/week, 3–27 MET-h/week, ≥27 MET-h/week), family history of colorectal cancer (no, yes), alcohol intake (<5 g/day, 5–15 g/day, ≥15 g/day), pack-years of smoking (0, 1–10, ≥10 pack-years), history of lower endoscopy (never/ever), caloric intake (quintiles), vitamin D (quintiles), calcium intake (quintiles), regular aspirin use (no, yes), folate intake (quintiles), MHT use (premenopausal, postmenopausal never user, postmenopausal past user, postmenopausal current user) (NHS and NHSII only) and red meat as main dish (quintiles).

bEvaluated in NHS and HPFS only.
has a long latency period of 10 or more years, and a long latency period was required to observe the protective effect of aspirin in prospective cohort studies.\textsuperscript{37} We considered several exposure windows, and observed no association with risk of colorectal cancer with medication use lagged up to 10 years. Furthermore, overall duration of follow-up in our study was up to 26 years, compared with up to 15 years in previous studies,\textsuperscript{26} additionally increasing our ability to detect any potential association.

To our knowledge, this is the first prospective cohort study to evaluate H2RA use in relation to risk of colorectal cancer. One previous prospective cohort study also found no overall increase in colorectal cancer risk in PPI users;\textsuperscript{37} however, this study had unclear time interval between exposure assessment and beginning of follow-up, lacked information on exposure and covariate updating, and did not consider some important confounders, such as history of colonoscopy. We were able to adjust for multiple confounders and examine multiple classifications of updated medication use, and we found no evidence of significant association between PPI use and risk of colorectal cancer.

Both PPI and H2RA lead to increased circulating levels of gastrin,\textsuperscript{7} which has been associated with increased risk of colorectal cancer in prospective studies.\textsuperscript{11} In animal models of colorectal cancer, gastrin leads to increased proliferation of colon mucosa and adenoma progression.\textsuperscript{8}–\textsuperscript{10} It is possible that the rise in gastrin levels due to PPI and H2RA use is not sufficient to promote development of colorectal tumours. Higher doses of medication over long periods of time may be necessary to exert an effect on tumour progression. While information on medication dosing was not available in our study, we observed no association with longer duration of use of acid-reducing medications with

| Table 4. Duration of acid-suppressive medications use and risk of colorectal cancer. |
|------------------------------------|----------------|----------------|----------------|----------------|
|                                   | No use | Used in 1–2 cycles | Used in ≥3 cycles | P-trend |
| PPI use                           |        |                   |                   |        |
| No. of cases                      | 952    | 159              | 58               |        |
| Person-years                      | 1,493,933 | 288,870         | 116,229          |        |
| Age-adjusted HR (95% CI)          | Ref.   | 0.80 (0.68–0.95) | 0.79 (0.60–1.04) | 0.01   |
| Multivariate HR (95% CI)\textsuperscript{a} | Ref.   | 0.89 (0.75–1.06) | 0.92 (0.69–1.21) | 0.21   |
| H2RA use                          |        |                   |                   |        |
| No. of cases                      | 1,821  | 225              | 94               |        |
| Person-years                      | 2,652,702 | 378,358         | 111,387          |        |
| Age-adjusted HR (95% CI)          | Ref.   | 0.84 (0.73–0.97) | 1.02 (0.82–1.25) | 0.33   |
| Multivariate HR (95% CI)\textsuperscript{a} | Ref.   | 0.91 (0.79–1.05) | 1.09 (0.88–1.35) | 0.95   |

\textsuperscript{a}Adjusted for age, BMI (< 25 kg/m\textsuperscript{2}, 25.1–27.5 kg/m\textsuperscript{2}, 27.6–30 kg/m\textsuperscript{2}, ≥30 kg/m\textsuperscript{2}), physical activity (<3 MET-h/week, 3–7 MET-h/week, ≥8 MET-h/week), family history of colorectal cancer (no, yes), alcohol intake (<5 g/day, 5–15 g/day, ≥15 g/day), pack-years of smoking (0, 1–10, ≥10 pack-years), history of lower endoscopy (never/ever), caloric intake (quintiles), vitamin D (quintiles), calcium intake (quintiles), regular aspirin use (no, yes), folate intake (quintiles), MHT use (premenopausal, postmenopausal never user, postmenopausal past user, postmenopausal current user) (NHS and NHSII only) and red meat as main dish (quintiles).

| Table 5. Association between colorectal cancer and lagged use of acid-suppressive medications. |
|------------------------------------|----------------|----------------|----------------|----------------|
|                                   | 2–4 years lag | 4–6 years lag | 6–8 years lag | 8–10 years lag |
|                                   | Non-users | Users | Non-users | Users | Non-users | Users | Non-users | Users | Non-users | Users |
| PPI use                           |           |       |           |       |           |       |           |       |           |       |
| No. of cases                      | 831      | 108   | 662       | 84    | 496       | 47    | 331       | 36    |           |       |
| Person-years                      | 1,373,645 | 219,447 | 1,118,996 | 163,824 | 877,318   | 115,076 | 631,274   | 69,835 |           |       |
| Age-adjusted HR (95% CI)          | Ref.     | 0.76 (0.62 to 0.94) | Ref. 0.81 (0.65 to 1.03) | Ref. 0.72 (0.53 to 0.97) | Ref. 1.01 (0.71 to 1.44) |
| Multivariate HR (95% CI)\textsuperscript{a} | Ref. 0.85 (0.69 to 1.05) | Ref. 0.90 (0.71 to 1.14) | Ref. 0.78 (0.58 to 1.07) | Ref. 1.12 (0.78 to 1.59) |
| H2RA use                          |           |       |           |       |           |       |           |       |           |       |
| No. of cases                      | 1,857    | 126   | 1,650    | 99    | 1,430     | 88    | 1,180     | 78    |           |       |
| Person-years                      | 2,615,912 | 179,320 | 2,274,028 | 153,533 | 1,954,486 | 131,335 | 1,639,395 | 109,876 |           |       |
| Age-adjusted HR (95% CI)          | Ref.     | 0.92 (0.76–1.10) | Ref. 0.82 (0.67–1.01) | Ref. 0.87 (0.70–1.08) | Ref. 0.96 (0.76–1.21) |
| Multivariate HR (95% CI)\textsuperscript{a} | Ref. 0.99 (0.83–1.19) | Ref. 0.89 (0.72–1.09) | Ref. 0.95 (0.76–1.18) | Ref. 1.02 (0.81–1.28) |

\textsuperscript{a}Adjusted for age, BMI (< 25 kg/m\textsuperscript{2}, 25.1–27.5 kg/m\textsuperscript{2}, 27.6–30 kg/m\textsuperscript{2}, ≥30 kg/m\textsuperscript{2}) physical activity (<3 MET-h/week, 3–7 MET-h/week, ≥8 MET-h/week), family history of colorectal cancer (no, yes), alcohol intake (<5 g/day, 5–15 g/day, ≥15 g/day), pack-years of smoking (0, 1–10, ≥10 pack-years), history of lower endoscopy (never/ever), caloric intake (quintiles), vitamin D (quintiles), calcium intake (quintiles), regular aspirin use (no, yes), folate intake (quintiles), MHT use (premenopausal, postmenopausal never user, postmenopausal past user, postmenopausal current user) (NHS and NHSII only) and red meat as main dish (quintiles).
 colorectal cancer. We also observed no increase in risk among patients with consistent PPI and H2RA use.

Furthermore, use of acid-suppressive medications leads to inhibition of gastric acid secretion and increased gastric pH, altering both the gastric microbiome and gut microbiome. However, it is not clear whether microbiome changes related to PPI use are overlapping with those associated with colorectal cancer. Our results do not suggest an adverse effect of microbiome changes caused by PPI use on colorectal cancer development.

A strength of this study is the prospective design that avoids the recall bias that often occurs in case–control studies, the large number of colorectal cancer cases and long follow-up in three independent cohorts of men and women. We collected information on the use of two acid-suppressive medications with different mechanisms of action; this use was updated every 2 years. Due to extensive information about lifestyle and dietary factors, we were able to adjust for important variables associated with both medication use and colorectal cancer, thereby minimising potential confounding. We were also able to evaluate confounding by indication, by accounting in our analyses for peptic or gastric ulcers, common reasons for use of acid-suppressive medications.

This study has several limitations. Since the questionnaire asked about “regular use” of the acid-suppressive medications in the previous 2-year cycle, without specifying the dose or frequency, we were not able to investigate a more precise dose–response relationship between acid-suppressive medication use and colorectal cancer. We had limited power to examine in more detail the associations with different tumour locations. Furthermore, evaluation of an even longer latency period might be necessary to further confirm the observed lack of association. Most of the participants in our cohort study were white, and our observations may therefore not be generalisable to other races and/or ethnicities.

In conclusion, in this prospective analysis among participants of three large cohorts, we observed no association between use of acid-suppressive medications and risk of colorectal cancer. Given the highly prevalent use of acid-suppressive medications in the US population, the lack of an association with increased colorectal cancer risk is reassuring.

ACKNOWLEDGEMENTS
We would like to thank the following state cancer registries for their help: AL, AZ, AR, CA, CO, CT, DE, FL, GA, ID, IL, IN, IA, KY, LA, ME, MD, MA, MI, NE, NH, NJ, NY, NC, ND, OH, OK, OR, PA, RI, SC, TN, TX, VA, WA and WY. The authors assume full responsibility for analyses and interpretation of these data.

AUTHOR CONTRIBUTIONS
Designed the study; A.B., K.N. Analysed the data: A.B., X.Z., K.N. Interpreted results: A.B., X.Z., V.M.-O., C.Y., A.T.C., S. 13. Imhann, F., Bonder, M. J., Vich Vila, A., Fu, J., Mujagic, Z., Vork, L. et al. Proton pump inhibitors affect the gut microbiome. Gut 65, 749–756 (2016).
14. Louis, P., Hold, G. L. & Flint, H. J. The gut microbiota, bacterial metabolites and colorectal cancer. Nat. Rev. Microbiol. 12, 661–672 (2014).
15. Sears, C. L. & Garrett, W. S. Microbes, microbiota, and colon cancer. Cell Host Microbe 15, 317–328 (2014).
16. Koritz, A. D., Gevers, D., Pedamallu, C. S., Michaud, M., Duke, F., Earl, A. M. et al. Genomic analysis identifies association of Fusobacterium with colorectal carcinoma. Genome Res. 22, 292–298 (2012).
17. Mima, K., Sukawa, Y., Nishihara, R., Qian, Z. R., Yamauchi, M., Inamura, K. et al. Fusobacterium nucleatum and T cells in colorectal carcinoma. JAMA Oncol. 1, 653–661 (2015).

Funding information This work was supported by following research grants from the National Institutes of Health (NIH: UM1 CA167552, U01 CA176726, U01 CA186107, P01 CA55075 and P01 CA87969. Additional support from NIH K07 CA222159 to A.B.; NIH K07 CA188126 and American Cancer Society Research Scholar Grant (RSG130476) to X.Z.; NIH R01 CA151993 and R35 CA197735 to S.O.; NIH R01 CA118553 to C.S.F.; Cancer Research UK Grand Challenge OPTIMISTIC Project award to S.O. and K.N.; Crohn’s and Colitis Foundation to P.L., NIH R01 CA205406, DOD CA163044 and Project P Fund to K.N. The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health. The funding sources played no role in study design, data collection, data analysis, interpretations of results or manuscript preparation.

Supplementary information is available for this paper at https://doi.org/10.1038/s41416-020-0939-y.

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REFERENCES
1. Center for Disease Control and Prevention. Selected Prescription Drug Classes Used in the Past 30 Days, By Sex and Age: United States, Selected Years 1988–1994 through 2011–2014 (2016).
2. Schubert, M. L. & Peura, D. A. Control of gastric acid secretion in health and disease. Gastroenterology 134, 1842–1860 (2008).
3. Savarino, V., Dulbecco, P., de Bortoli, N., Ottone, A. & Savarino, E. The appropriate use of proton pump inhibitors (PPIs): need for a reappaisal. Eur. J. Intern. Med. 37, 19–24 (2017).
4. Johnson, D. A., Katz, P. O., Armstrong, D., Cohen, H., Delaney, B. C., Howden, C. W. et al. The safety of appropriate use of over-the-counter proton pump inhibitors: an evidence-based review and Delphi consensus. Drugs 77, 547–561 (2017).
5. Festa, H. P., Thijs, J. C., Lamers, C. B., Jansen, J. M., Pols, G., Frants, R. et al. Effect of oral omeprazole on serum gastrin and serum pepsinogen I levels. Gastroenterology 87, 1030–1034 (1984).
6. Lazzaroni, M., Sangalotti, O. & Bianchi Porro, G. Gastric acid secretion and plasma gastrin during short-term treatment with omeprazole and ranitidine in duodenal ulcer patients. Hепato-Gastroenterology 39, 366–370 (1992).
7. Ohsawa, T., Hirata, W. & Higuchi, S. Effects of three H2-receptor antagonists (cimetidine, famotidine, ranitidine) on serum gastrin level. Int. J. Clin. Pharmac. Res. 22, 29–35 (2002).
8. Koh, T. J., Dockray, G. J., Varro, A., Cahlil, R. J., Dangler, C. A., Fox, J. G. et al. Overexpression of glycine-extended gastrin in transgenic mice results in increased colonic proliferation. J. Clin. Invest. 103, 1119–1126 (1999).
9. Watson, S. A., Morris, T. M., McWilliams, D. F., Harris, J., Evans, S., Smith, A. et al. Potential role of endocrine gastrin in the colonic adenoma carcinoma sequence. Br. J. Cancer 87, 567–573 (2002).
10. Watson, S. A. & Smith, A. M. Hypergastrinemia promotes adenoma progression in the APC(Min−/−) mouse model of familial adenomatous polyposis. Cancer Res. 61, 625–631 (2001).
11. Thorburn, C. M., Friedman, G. D., Dickinson, C. J., Vogelman, J. H., Orentreich, N. & Parsonnet, J. Gastrin and colorectal cancer: a prospective study. Gastroenterology 115, 275–280 (1998).
12. Freedberg, D. E., Lebowohl, B. & Abrams, J. A. The impact of proton pump inhibitors on the human gastrointestinal microbiome. Clin. Lab. Med. 34, 771–785 (2014).
13. Imhann, F., Bonder, M. J., Vich Vila, A., Fu, J., Mujagic, Z., Vork, L. et al. Proton pump inhibitors affect the gut microbiome. Gut 65, 740–748 (2016).
14. Jackson, M. A., Goodrich, J. K., Maxan, M. E., Freedberg, D. E., Abrams, J. A., Poole, A. C. et al. Proton pump inhibitors alter the composition of the gut microbiota. Gut 65, 749–756 (2016).
15. Louis, P., Hold, G. L. & Flint, H. J. The gut microbiota, bacterial metabolites and colorectal cancer. Nat. Rev. Microbiol. 12, 661–672 (2014).
16. Sears, C. L. & Garrett, W. S. Microbes, microbiota, and colon cancer. Cell Host Microbe 15, 317–328 (2014).
17. Koritz, A. D., Gevers, D., Pedamallu, C. S., Michaud, M., Duke, F., Earl, A. M. et al. Genomic analysis identifies association of Fusobacterium with colorectal carcinoma. Genome Res. 22, 292–298 (2012).
18. Mima, K., Sukawa, Y., Nishihara, R., Qian, Z. R., Yamauchi, M., Inamura, K. et al. Fusobacterium nucleatum and T cells in colorectal carcinoma. JAMA Oncol. 1, 653–661 (2015).
19. Nguyen, P. T., Baldeck, J. D., Olsson, J. & Marquis, R. E. Antimicrobial actions of benzimidazoles against oral streptococci. *Oral Microbiol. Immunol.* **20**, 93–100 (2005).

20. Sheng, J., Nguyen, P. T., Baldeck, J. D., Olsson, J. & Marquis, R. E. Antimicrobial actions of benzimidazoles against the oral anaerobes *Fusobacterium nucleatum* and *Prevotella intermedia*. *Arch. Oral Biol.* **51**, 1015–1023 (2006).

21. Chubak, J., Boudreau, D. M., Rulyak, S. J. & Mandelson, M. T. Colorectal cancer risk in relation to use of acid suppressive medications. *Pharmacoepidemiol. Drug Saf.* **18**, 540–544 (2009).

22. Lai, S. W., Liao, K. F., Lai, H. C., Lin, C. L. & Sung, F. C. Use of proton pump inhibitors correlates with increased risk of colorectal cancer in Taiwan. *Asia Pac. J. Clin. Oncol.* **9**, 192–193 (2013).

23. Robertson, D. J., Larsson, H., Friis, S., Pedersen, L., Baron, J. A. & Sorensen, H. T. Proton pump inhibitor use and risk of colorectal cancer: a population-based, case–control study. *Gastroenterology* **133**, 755–760 (2007).

24. Siersma, P. D., Yu, S., Sahbaie, P., Steyerberg, E. W., Simpson, P. W., Kuipers, E. J. et al. Colorectal neoplasia in veterans is associated with Barrett’s esophagus but not with proton-pump inhibitor or aspirin/NSAID use. *Gastrointest. Endosc.* **63**, 581–586 (2006).

25. van Soest, E. M., van Rossum, L. G., Dieleman, J. P., van Oijen, M. G., Siersema, P. D., Sturkenboom, M. C. et al. Proton pump inhibitors and the risk of colorectal cancer. *Am. J. Public Health* **106**, 1599–1607 (2016).

26. Krohn, K. M., Hennekens, C. H., Rosner, B. & Speizer, F. E. The nurses’ health study. *Am. J. Nurs.* **78**, 1039–1040 (1978).

27. Colditz, G. A. & Hankinson, S. E. The Nurses’ Health Study: lifestyle and health among women. *Nat. Rev. Cancer* **5**, 388–396 (2005).

28. Troy, L. M., Michels, K. B., Hunter, D. J., Spiegelman, D., Manson, J. E., Colditz, G. A. et al. Self-reported birthweight and history of having been breastfed among younger women: an assessment of validity. *Int. J. Epidemiol.* **25**, 122–127 (1996).

29. Giovannucci, E., Ascherio, A., Rimm, E. B., Colditz, G. A., Stampfer, M. J. & Willett, W. C. Physical activity, obesity, and risk for colon cancer and adenoma in men. *Ann. Intern. Med.* **122**, 327–334 (1995).

30. Lee, D. H., Keum, N. & Giovannucci, E. L. Colorectal cancer epidemiology in the Nurses’ Health Study. *Am. J. Public Health* **106**, 1599–1607 (2016).

31. Bardou, M., Barkun, A. N. & Martel, M. Obesity and colorectal cancer. *Gut* **62**, 933–947 (2013).

32. Zauber, A. G., Winawer, S. J., O’Brien, M. J., Landsdorp-Vogelaar, I., van Ballegooijen, M., Hankey, B. F. et al. Colonoscopic polypectomy and long-term prevention of colorectal-cancer deaths. *N. Engl. J. Med.* **366**, 687–696 (2012).

33. Giovannucci, E., Egan, K. M., Hunter, D. J., Stampfer, M. J., Colditz, G. A., Willett, W. C. et al. Aspirin and the risk of colorectal cancer in women. *N. Engl. J. Med.* **333**, 609–614 (1995).

34. Minalyan, A., Gabrielyan, L., Scott, D., Jacobs, J. & Pisegna, J. R. The gastric and intestinal microbiome: role of proton pump inhibitors. *Curr. Gastroenterol. Rep.* **19**, 42 (2017).