Short-Term Acetylsalicylic Acid (Aspirin) Use for Pain, Fever, or Colds – Gastrointestinal Adverse Effects
A Meta-Analysis of Randomized Clinical Trials

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Abstract

Background and Aim: Acetylsalicylic acid (ASA [aspirin]) is a commonly used over-the-counter drug for the treatment of pain, fever, or colds, but data on the safety of this use are very limited. The aim of this study was to provide data on the safety of this treatment pattern, which is of interest to clinicians, regulators, and the public.

Methods: A meta-analysis of individual patient data from 67 studies sponsored by Bayer HealthCare was completed. The primary endpoints were patient-reported gastrointestinal (GI) adverse events (AEs); the secondary endpoints were the incidence of patient-reported non-GI AEs. Event incidence and odds ratios (ORs) based on Cochran-Mantel-Haenszel estimates are reported. In total, 6181 patients were treated with ASA, 3515 with placebo, 1145 with acetaminophen (paracetamol), and 754 with ibuprofen. Exposure to ASA was short term (82.5% of patients had a single dose).

Results: GI AEs were more frequent with ASA (9.9%) than with placebo (9.0%) [OR 1.3; 95% CI 1.1, 1.5]. Dyspeptic symptoms were infrequent (4.6% in placebo subjects). The ORs for ASA were 1.3 (95% CI 1.1, 1.6) versus placebo; 1.55 (95% CI 0.7, 3.3) versus ibuprofen; and 1.04 (95% CI 0.8, 1.4) versus acetaminophen. There were very few serious GI AEs (one ASA case; three placebo cases). No differences were found for non-GI AEs and no cases of cerebral hemorrhage were reported.
Conclusion: Short-term, mostly single-dose exposure to ASA for the treatment of pain, fever, or colds was associated with a small but significant increase in the risk of dyspepsia relative to placebo. No serious GI complications were reported.

Background

Acetylsalicylic acid (ASA [aspirin]) at doses of ≤325 mg/day is used for the treatment and prevention of cardiovascular events.[1] At higher doses, used for the treatment of pain, fever, or colds, it is one of the most commonly used over-the-counter (OTC) drugs worldwide.[2] The recommended OTC dose for these indications varies by country, but is generally 500–1000 mg as a single dose and 3000–4000 mg/day. Marketing surveys indicate that treatment is usually short term for acute pain, suggesting that most people take one to two tablets for 1 day.[3]

At these doses, ASA can be considered a traditional nonselective NSAID.[4] These compounds are associated with an increased risk of side effects, and those in the gastrointestinal (GI) tract are among the most frequently reported.[5] GI complications are life-threatening events associated with NSAID use, but other adverse effects such as dyspepsia are also important since they may lead to avoidance of treatment.[5,6]

There are only limited data on GI safety regarding short-term ASA use for the treatment of various acute conditions. Observational studies have reported the relative risks of upper GI bleeding associated with ASA above 500 mg/day to be similar to those of other NSAIDs, but information on dose, duration of treatment, type of use, and indication is often limited or absent. Observational studies often do not differentiate between chronic and acute OTC ASA use or they do not capture OTC use at all, severely limiting the interpretation of the data.[7-9] Moreover, data regarding more frequent GI side effects associated with NSAID use such as dyspepsia are rarely reported in the literature.[10] Data on doses of ASA used for the treatment of pain, fever, or colds have been even less frequently reported.

Other side effects associated with ASA use include intracranial bleeding, other non-GI bleeding, tinnitus, dizziness, headache, impaired hearing, hypersensitivity reactions, and mental confusion.[11] Data on the occurrence of these side effects during short-term ASA use for acute conditions have rarely been reported.

To investigate these adverse events (AEs), we evaluated the safety profile of short-term ASA use at the recommended doses for various OTC ASA indications, based on individual subject data obtained from all clinical trials with doses of ASA ≥325 mg/day conducted by Bayer HealthCare between 1987 and 2008.

Methods

Setting

Individual patient data for the meta-analysis were obtained from all studies conducted by Bayer HealthCare by 31 March 2008, where ASA was evaluated in a clinical trial setting and where adequate data documentation in terms of AE reporting was available. The data pool contained 87 studies in total. Among these, 33 were double-blind, 2 were single-blind, 31 were open-label, and the blinding in one study was not recorded. All studies included were either efficacy or pharmacokinetic studies. Studies of low-dose ASA for the prevention of cardiovascular diseases (daily dose ≤325 mg) were excluded. As a result, data from 67 studies were considered for this report.

The most relevant inclusion criteria in these studies were as follows: (i) patient presented with the investigated indication in pain, fever, or colds, and was otherwise healthy (efficacy study); and (ii) volunteers (pharmacokinetic studies). The most relevant exclusion criteria were as follows: (i) history of presence of asthma or hypersensitivity to ASA, salicylate, or NSAIDs; (ii) active peptic ulcer; (iii) history of gastroduodenal bleeding; (iv) hemorrhagic diathesis; (v) impaired hepatic function; or (vi) impaired renal function.
Endpoints

The primary endpoints were patient-reported GI AEs. During the studies, trial subjects were asked to report any AE, investigators were instructed to provide clinical diagnosis of AEs when possible, and Bayer HealthCare assigned the appropriate Medical Dictionary for Regulatory Activities (MedDRA) term to each AE. MedDRA codes (version 11.0) were used for the identification of AEs of interest in all clinical trials in the database, either according to predefined, standardized MedDRA queries or according to selected MedDRA preferred terms, high-level terms, high-level group terms, or system organ classes. The four academic authors (AL, DM, SS, JB) defined a priori the choice of terms for appropriate retrieval of these events; the same authors also defined the additional events of interest for the so-called combined preferred terms per system organ class. GI preferred terms were combined into the following variables: GI bleeding, dyspepsia, severe dyspepsia, minor dyspepsia, severe dyspepsia, any dyspepsia, abdominal pain, GI bleeding, gastroesophageal reflux disease (GERD)-related symptoms. Definitions of these combined preferred terms are included in table I.

Secondary endpoints were patient-reported non-GI AEs, including cerebral hemorrhage, other bleeding (non-GI, non-cerebral), hypersensitivity reactions, headache, dizziness, impaired hearing ability, tinnitus, mental confusion, oral complications and ‘signs of overdose’ as a composite of the following AEs of interest: headache, dizziness, impaired hearing ability, tinnitus, and mental confusion. MedDRA codes were also used for identification of these AEs of interest. For both primary and secondary endpoints, a time window of 7 days after drug discontinuation was used to include events from any treatment arm that were defined as ‘treatment related.’

Treatments Considered for the Analysis

Clinical trials in the Bayer HealthCare database that studied treatment with ASA alone and in combination with another active ingredient (pseudoephedrine, lidocaine, or dextromethorphan) versus placebo or active comparator. For the purpose of this analysis, combinations of ASA plus vitamin C, ASA plus caffeine, and ASA plus calcium were considered to be ‘ASA alone.’ This study reports on comparisons between two different treatment arms: ASA versus placebo, each given alone or each combined with additional therapy (e.g. pseudoephedrine, lidocaine, or dextromethorphan). In addition, some studies were an ‘active comparator.’ These were acetaminophen (paracetamol), carried out comparing ASA with ibuprofen, pseudoephedrine, sumatriptan, lidocaine, ketoprofen, or, in a few cases, ergotamine tartrate and ASA-acetaminophen-caffeine combinations. We report events from

| Combined PT | Included PTs |
|-------------|-------------|
| Dyspepsia   | Dyspepsia   |
|             | Epigastric discomfort |
|             | Eructation    |
| Minor dyspepsia | Abdominal discomfort |
|             | Dyspepsia   |
|             | Epigastric discomfort |
|             | Eructation    |
|             | Flatulence   |
|             | Gastric dilatation |
|             | Gastric disorder |
|             | Hyperchlorhydria |
|             | Nausea       |
|             | Stomach discomfort |
|             | Abdominal pain upper |
| Severe dyspepsia | Retching |
|             | Vomiting     |
| Any dyspepsia | Included PTs for ‘Minor dyspepsia’ and ‘Severe dyspepsia’ combined |
| Abdominal pain | Abdominal pain |
|             | Abdominal pain lower |
| GI bleeding  | Haematemesis |
|             | Haematochezia |
|             | Melaena      |
| GERD-related symptoms | Gastrooesophageal reflux disease |
|             | Dysphagia    |
|             | Oesophageal pain |
|             | Salivary hypersecretion |

GERD = gastroesophageal reflux disease.
studies comparing ASA with either acetaminophen or ibuprofen, since only clinical trials with these compounds are of major clinical interest and only these studies had a sufficient number of cases to carry out appropriate analyses.

**Data Extraction and Management**

Data management and statistical evaluation was performed using the SAS® software package version 9.2 (SAS Institute Inc., Cary, NC, USA). The database structure was developed between Bayer HealthCare and M.A.R.C.O., Dusseldorf, Germany, an independent institute for clinical research and statistics. Data were provided by Bayer HealthCare in one of the following formats: data listings derived from medical research reports on paper, data listings from medical research reports in PDF, or SAS® datasets. Data listings provided as paper copies were entered into the database, either by independent double-data entry with subsequent comparison of the datasets and entry-error correction, or, in case of small studies with few data, by single data entry with a subsequent 100% visual inspection. Data listings provided in PDF were copied into Microsoft® Excel (Microsoft Corporation, Redmond, WA, USA) and then imported into SAS®. The SAS® datasets were then transformed into the target database structure using SAS® modification programs. Information derived from the medical research reports concerning study title, design, blinding, randomization, dosing, and so forth was integrated into the target database. Appropriate quality control checks were in place at each data management step.

**Statistics**

The scope of the analysis, statistical methods, and content of tables and graphs was documented and pre-specified in a statistical analysis plan (SAP) by all authors prior to start of the analysis. We have estimated incidence rates for both AEs overall and those events specified by the trial investigators as adverse drug related (i.e. adverse drug reactions [ADRs]). All statistical analyses were descriptive in nature. Incidence rates calculated for this meta-analysis refer to the number of subjects who reported at least one event in the numerator and the number of all subjects under observation in the denominator.

**Measures of Treatment Effect**

Analysis of risk differences and odds ratios (ORs) were performed for all events of interest and treatment comparisons defined in the Treatments Considered for the Analysis section. In all analyses, trial was used as a stratum, as is standard in meta-analysis.[12,13] These analyses were performed twice, excluding and including studies with no AEs (‘zero-event studies’) in both treatment arms and for all reported AEs, as well as drug-related AEs, yielding a total of 1280 analyses. Confidence interval plots (forest plots) and radial plots (Galbraith plots) were provided for the risk analyses. Estimated ORs and risk differences were based on Cochran-Mantel-Haenszel estimates, as this is robust even in ‘sparse data’ stratifications (i.e. where few cases of AEs occur). Weighted estimates for ORs and risk differences were calculated[14-16] and reported. For handling of zero-event studies, a standardized continuity correction of 0.05 was used, taking into account the size of treatment arms.[16] This was a deviation from the SAP, where a factor of 1/10000 was proposed.

For analysis of heterogeneity over studies, a modified Breslow-Day statistic was used in the OR analyses[17,18] and the Cochran-Mantel-Haenszel Q-statistic was used in the risk difference analyses.[19] A p-value of 0.10 was considered to indicate heterogeneity.

Assessment of ASA-by-subgroup interactions was carried out for sex and age. For each event of interest, a Cochran-Mantel-Haenszel risk analysis was performed and the OR for the comparisons was estimated separately between male and female subjects. The two estimates were then compared and the difference calculated (log OR [ASA vs placebo for male patients] minus log OR [ASA vs placebo for female patients]). The variance of this contrast was calculated as the sum of the variances of the two log ORs. The square root of this was taken as the standard error and used to calculate 95% confidence intervals. The test statistic for interaction was the ratio of this difference to its standard error.[20] Analogously, estimates for the ASA-by-age interaction were derived.
Two different subgroup allocations were considered: patients <40 versus patients ≥40 years of age and patients <65 versus patients ≥65 years of age. Subgroup analyses concerning daily dose, treatment duration, and treatment formulation were also performed. A p-value of <0.10 was assumed to provide evidence of heterogeneity.

Both treatment arms were included for parallel-group studies; only data from the first period were included for crossover studies. Descriptive, exploratory evaluations, and statistical analyses were based on available data. AEs with incomplete or missing onset dates were considered as treatment related. The same approach was also used if no dosing date was given.

All patients treated at least once were included in the analysis. For the subgroup analyses, patients were grouped according to combinations of treatment duration (single dose, multiple dose, all) and formulation (plain: tablets, coated tablets, capsules, dry granules; effervescent tablets; intravenous; all). Multiple dosing was defined as more than one dose of study drug in any of the treatment groups.

**Results**

**Description of Trials and Demographics of Patients**

A total of 67 studies were included in the analysis (figure 1). The aim of the majority of the studies included was the evaluation of the efficacy of ASA for the treatment of pain, fever, or colds; a small number were conducted to evaluate the pharmacokinetics of ASA. Overall, exposure to
ASA and its combinations was brief. Of the 6181 patients allocated to ASA and ASA combinations, 5099 (82.5%) received a single dose, 1082 (17.5%) received a multiple-dose regimen, and 188 (3.0%) were treated for more than 5 days. Over half (54%; 3337 patients) of the ASA study population took a daily dose between 500 and 1000 mg (figure 2).

The daily doses of acetaminophen taken by patients in these trials were 300 mg (31%), 500 mg (16.6%), and 1000 mg (52.3%), whereas the doses of ibuprofen were 200 mg (45%) and 400 mg (55%). The demographics of patients are included in table II.

**Safety Analysis**

The overall incidence of AEs was low, and was similar between subjects treated with ASA (741/4884; 15.2%) and those treated with placebo (580/3731; 15.5%) [OR 1.1; 95% CI 0.96, 1.2]; risk difference 1.0% (95% CI –0.5, 2.5). GI AEs of any severity were more frequent in the ASA groups than in the placebo groups, as summarized in table III. Risks of abdominal pain and GERD-related symptoms were non-significantly increased in the ASA groups; severe dyspepsia was non-significantly less common in ASA subjects. There were very few serious GI adverse events; one case of acute appendicitis in the ASA plus pseudoephedrine arm and two cases in the placebo arm (one hematemesis and one appendicitis perforation). Neither the appendicitis nor the appendicitis perforation events were considered drug related by the original study investigators. GI hemorrhage (hematemesis) was reported once among subjects randomized to ASA plus dextromethorphan and three times among subjects randomized to placebo (hematemesis, hematochezia, and melela). Similar figures were observed when events were analyzed as drug-related AEs, although absolute numbers were smaller (figure 3).

No cerebral hemorrhage or hypersensitivity reaction occurred in any of the trials, although one occurrence of impaired hearing ability and another of mental confusion were reported in the ASA groups. Other non-GI AEs are also summarized in table III. There was a reduction in the risk of headache from ASA and there were non-significant increases in tinnitus and non-GI bleeding. Very similar results to those reported in table III were obtained when ASA in monotherapy was compared with placebo alone (data not shown).[21]

There was no indication of heterogeneity across studies in AEs for the OR analysis; there was statistically significant heterogeneity in the risk difference analysis for the terms GI AE, dyspepsia, minor dyspepsia, any dyspepsia, headache, signs of overdose, and oral complications. Since
ORs were considered to be the main parameter, no steps were undertaken to correct for statistical heterogeneity in this analysis of pooled data.

Additional analyses were made according to ASA formulation, median daily ASA dose, and single- versus multiple-dose regimens. Overall, compared with plain and effervescent oral formulations, the incidence of AEs with exposure to intravenous ASA was lower (data not shown).[21] More patients receiving doses >1000 mg/day reported AEs (18.1%) than those receiving doses <500 mg/day (15.1%) or those receiving placebo (13.5%). Multiple-dose administration of ASA (up to 5 days) was associated with an increased incidence of AEs (16.2%) when compared with single-dose administration (12.8%), and this was similar to results observed in the placebo arm (17.0% and 12.8%, respectively).

Acetylsalicylic acid (ASA (Aspirin))-by-Subgroup Interaction

Analyses for detection of an interaction between age (<65 vs ≥65 years of age; <40 vs ≥40 years of age) or sex (male vs female) and the risk of experiencing an AE were carried out for the comparison of ASA versus placebo. There was no interaction by age or sex for any of the events of interest. For the combined term ‘minor dyspepsia’, a borderline statistically significant ($p=0.052$) interaction effect was observed for patients ≥40 years of age when compared with those <40 years of age (OR 1.98 [95% CI 1.2, 3.1]). Approximately 1% of the patients were ≥65 years old (1.1% of patients on ASA, 1.3% of patients on placebo) and therefore no meaningful data can be reported for these subgroups.

| Table II. Characteristics of subjects included in the meta-analysis |
|---------------------------|---------------------------|---------------------------|---------------------------|
| Parameter                  | Placebo                        | All acetylsalicylic acid (aspirin) treatments | Active comparator | Total                |
| Sex [n (%)]                |                                |                                |                |
| Men                        | 1257 (36)                      | 2638 (43)                      | 1260 (36)      | 5155 (39)           |
| Women                      | 2258 (64)                      | 3543 (57)                      | 2266 (64)      | 8067 (61)           |
| Race/ethnicity [n (%)]     |                                |                                |                |
| Unknown                    | 314 (9)                        | 916 (15)                       | 955 (27)       | 2185 (17)           |
| Caucasian                  | 2917 (83)                      | 4843 (78)                      | 2351 (67)      | 10 111 (76)         |
| Black                      | 146 (4)                        | 217 (4)                        | 103 (3)        | 466 (4)             |
| Asian                      | 27 (1)                         | 41 (1)                         | 20 (1)         | 88 (1)              |
| Hispanic                   | 74 (2)                         | 93 (2)                         | 51 (1)         | 218 (2)             |
| Other                      | 37 (1)                         | 71 (1)                         | 46 (1)         | 154 (1)             |
| Age [y]                    |                                |                                |                |
| n                          | 3465                           | 6131                           | 3276           | 12 872              |
| Mean                       | 33.22                          | 32.51                          | 33.30          | 32.9                |
| SD                         | 12.92                          | 12.67                          | 13.28          | 12.9                |
| Range                      | 15–77                          | 15–81                          | 15–78          | 15–81               |
| Weight [kg]                |                                |                                |                |
| n                          | 3463                           | 6088                           | 3275           | 12 826              |
| Mean                       | 71.31                          | 72.31                          | 72.16          | 72.00               |
| SD                         | 14.98                          | 15.14                          | 15.29          | 15.14               |
| Range                      | 40–158                         | 35–167                         | 37–159         | 35–167              |
| BMI [kg/m²]                |                                |                                |                |
| n                          | 3463                           | 6088                           | 3274           | 12 825              |
| Mean                       | 24.65                          | 24.71                          | 24.90          | 24.74               |
| SD                         | 4.50                           | 4.58                           | 4.64           | 4.57                |
| Range                      | 12.5–56.2                      | 14.5–60.6                       | 15.4–60.1      | 12.5–60.6           |

BMI = body mass index.

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ASA versus Active Comparator

This analysis was carried out comparing ASA monotherapy (all doses) with acetaminophen (300–1000 mg/day) or ibuprofen (200–400 mg/day). Table IV summarizes the rates and ORs for the different GI and non-GI AEs. In general, the rates were low for all AEs. No statistically significant differences for the combined terms of dyspepsia were found between treatments, although the ORs were >1.0 for the comparisons with ibuprofen. Similar figures were observed when data were reported as ADRs (data not shown).[21]

Discussion

This meta-analysis has shown that the risk of AEs from ASA was low at the doses and durations commonly consumed as OTC medication for the relief of pain, fever, or colds. In particular, there was a virtual absence of major GI (e.g. bleeding or perforation) and non-GI complications (e.g. cerebral hemorrhage). The use of ASA was associated with a small increase of the risk of dyspepsia when compared with placebo and with a non-significant increase in comparison with ibuprofen, another OTC drug also commonly used for the treatment of colds or acute pain. There was an increased risk of dyspepsia with ASA versus acetaminophen when considering the ‘dyspepsia’ MedDRA term, but not when the broader, combined dyspepsia terms defined a priori by the academic authors were used (table I).

It is important to emphasize that the conclusions obtained from this study on the safety of this therapeutic approach with ASA apply only to very short-term treatment. The majority of patients included in these trials took a single dose of 500–1000 mg in 1 day and were relatively young. Nonetheless, the data reported are clinically valuable. First, previous information on the safety of ASA at OTC doses has relied on a small number of observational studies, since many investigations do not report on OTC use. Second, the

Table III. Gastrointestinal (GI) adverse events (AEs) and non-GI AEs occurring in patients treated with acetylsalicylic acid (ASA [aspirin]) or placebo

| Event                                | ASA arm [%] (n = 4884) | Placebo arm [%] (n = 3731) | OR (95% CI) | Risk difference [%] (95% CI) |
|--------------------------------------|------------------------|-----------------------------|-------------|-----------------------------|
| **GI AEs**                           |                        |                             |             |                             |
| All GI AEs                           | 9.9                    | 9.0                         | 1.3 (1.1, 1.5) | 2.1 (0.9, 3.3) |
| dyspepsiaa                           | 1.8                    | 1.4                         | 1.7 (1.2, 2.4) | 0.8 (0.2, 1.3) |
| Minor GI disordersb                  | 5.0                    | 4.5                         | 1.2 (1.0, 1.5) | 0.9 (0.01, 1.8) |
| abdominal painc                      | 0.5                    | 0.2                         | 1.9 (0.9, 4.0) | 0.2 (0.0, 0.5) |
| any dyspepsia                       | 5.3                    | 4.6                         | 1.3 (1.1, 1.6) | 1.2 (0.3, 2.2) |
| minor dyspepsia                      | 5.0                    | 4.0                         | 1.4 (1.1, 1.8) | 1.5 (0.6, 2.3) |
| severe dyspepsia                      | 0.6                    | 0.9                         | 0.7 (0.4, 1.2) | –0.3 (–0.6, 0.1) |
| GERD-related symptomsd               | <0.1                   | <0.1                        | 1.5 (0.3, 7.0) | 0.03 (–0.09, 0.1) |
| **Non-GI AEs**                      |                        |                             |             |                             |
| Headache                             | 0.8                    | 1.7                         | 0.5 (0.3, 0.8) | –0.7 (–1.2, –0.2) |
| Dizziness                            | 0.9                    | 1.1                         | 0.9 (0.6, 1.3) | –0.1 (–0.6, 0.3) |
| Tinnitus                             | 0.2                    | 0.1                         | 1.7 (0.6, 4.5) | 0.1 (–0.1, 0.3) |
| Sign of overdose                     | 1.9                    | 2.8                         | 0.7 (0.6, 1.0) | –0.6 (–1.3, 0.0) |
| Other bleeding (non-GI, non cerebral) | 0.3                    | 0.2                         | 1.5 (0.6, 3.4) | 0.1 (–0.1, 0.3) |
| Oral complicationsd                  | 2.9                    | 3.0                         | 1.3 (0.97, 1.7) | 0.6 (–0.1, 1.3) |

a MedDRA term.
b Minor GI disorders include heartburn, nausea, vomiting and abdominal pain. Other terms are defined in table I.
c Combined term.
ASA arm = ASA alone or combined with additional therapy; GERD = gastroesophageal reflux disease; MedDRA = Medical Dictionary for Regulatory Activities; OR = odds ratio; Placebo arm = placebo alone or combined with the additional therapy (pseudoephedrine, lidocaine, or dextromethorphan).
data we report here are valid for typical OTC ASA use in the general population. According to some surveys on consumer use of analgesics,\(^3,22\) 92.4% of ASA users take one to two tablets (500 mg tablet equivalent) in a 4-week period.

Our meta-analysis agrees with two other studies reported by Edwards et al.,\(^23\) and Steiner and Voelker.\(^24\) The former study was not a pooled analysis of individual patient data and was just focused on the efficacy of single-dose ASA for the relief of postoperative pain. Nevertheless, they found risks of AEs with ASA use (13%) and placebo (11%) similar to those reported here. They also found a significantly higher incidence of drowsiness and what they describe as ‘gastric irritation,’ with ASA 600–650 mg than with placebo. The latter study\(^24\) used pooled individual patient data from 1581 patients treated with ASA and 1271 patients treated with placebo in nine randomized, double-blind, placebo-controlled clinical trials, and these were included in our study. They were chosen because all patients used single doses of ASA 1000 mg for the treatment of acute migraine attacks, episodic tension-type headache, and dental pain. The reported AE rates for ASA and placebo were 14.9% and 11.1%, respectively; this included 5.9% of patients experiencing GI events with ASA and 3.5% experiencing them with placebo. One additional study\(^11\) reported side effects arising from ASA, acetaminophen, codeine, or placebo therapy in a retrospective analysis of 54 single-dose, double-blind studies involving 4346 patients with postsurgical dental pain. Both the overall and the GI AEs were low for all treatments, but codeine was associated with a higher incidence of nausea, drowsiness, and dizziness than acetaminophen, ASA, or placebo.

Available observational studies reporting on the GI safety of non-cardiovascular ASA, taken for the treatment of pain, fever, or colds, indicate that its use is associated with an increased risk of major GI complications, similar to other non-selective NSAIDs.\(^6-8\) In contrast, in our data and in those of Edwards et al.,\(^23\) no serious GI complications were associated with ASA use or active comparators. This should not necessarily be seen as contradictory, since it is reasonable to assume that the populations and type of drug use studied are different. Clinical trials usually exclude patients with known GI risk factors, whereas observational studies include patients with various levels of risk. This meta-analysis probably included a population that was healthier and younger than that reported in observational studies. Furthermore, it is possible that observational studies report on patients with longer exposure to ASA use, whereas our study basically applies to those taking a single tablet or using ASA doses no higher

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**Fig. 3.** Frequencies of patients who reported at least one adverse event (AE) of interest (all and drug related) with acetylsalicylic acid (ASA [aspirin]) or with placebo. ADR = adverse drug reaction; GI = gastrointestinal; PT = preferred term; SOC = system organ class.
than 1000 mg/day for the relief of pain, fever, or colds. The same considerations may apply to the absence of GI bleeding events with ibuprofen in this study, where the doses used were low (200–400 mg/day).

ASA use has been associated with other non-GI AEs, both serious and less serious.\cite{11,25,26} No cases of serious non-GI AEs were reported and no significant differences were found between ASA and placebo in this meta-analysis.

We did not find statistically significant heterogeneity in the ORs in our analysis, suggesting that these statistics are probably generalizable, at least to the sort of patient included in the studies. However, as might be expected with relatively constant ORs and differing background rates of AEs, there was heterogeneity in the risk differences. This finding suggests that the risk differences cannot be directly applied in other clinical contexts.

The data presented in this study have several limitations. The way ASA is used may vary among different populations and therefore the data reported may not be applicable to them. The report from Curhan et al.\cite{2} from the Nurses’ Health Study showed that ASA was used daily by 25.4% of women >51 years of age, and that 40.1% used ASA >1 day/week. Since it is now widely known that ASA protects against cardiovascular events and colon cancer,\cite{1,27} some may take this compound very often or on a daily basis for these purposes. In any case, multiple-day exposure to ASA is not uncommon in clinical practice, and the data provided here, although focused on short-term use of ASA for the treatment of acute pain, fever, or colds, suggest an increased risk of AEs with increasing dose and multiple doses.

Another limitation of this study is that the safety data reported here may not be valid for the entire population with access to OTC ASA for the relief of pain, fever, or colds. The risk factors for GI complications in NSAID and ASA users are well defined and include older age (>65 years old), patients with ulcer history, and patients with concomitant use of other NSAIDs, corticosteroids,

Table IV. Adverse events (AEs) with acetylsalicylic acid (ASA [aspirin]) in monotherapy compared with active comparators (acetaminophen [paracetamol] and ibuprofen)

| Event                        | ASA [%] (n = 1128) | Acetaminophen [%] (n = 1145) | OR (95% CI)      | ASA [%] (n = 707) | Ibuprofen [%] (n = 754) | OR (95% CI)      |
|------------------------------|--------------------|-------------------------------|------------------|--------------------|------------------------|------------------|
| GI AEs                       | 10.4               | 10.1                          | 1.0 (0.8, 1.4)   | 3.5                | 2.3                    | 1.4 (0.8, 2.7)   |
| dyspepsia\(^a\)              | 4.0                | 2.5                           | 1.7 (1.0, 2.8)   | 0.1                | 0.4                    | 0.3 (0.0, 2.9)   |
| Minor GI disorders\(^b\)     | 6.8                | 7.2                           | 0.9 (0.7, 1.3)   | 3.4                | 1.9                    | 1.7 (0.9, 3.3)   |
| abdominal pain\(^c\)         | 0.3                | <0.1                          | 2.5 (0.3, 18.7)  | 0.3                | 0.3                    | 1.0 (0.1, 6.4)   |
| any dyspepsia\(^c\)          | 7.9                | 7.6                           | 1.0 (0.7, 1.4)   | 2.7                | 1.6                    | 1.5 (0.7, 3.2)   |
| minor dyspepsia\(^c\)        | 7.6                | 7.2                           | 1.1 (0.8, 1.5)   | 2.5                | 1.3                    | 1.8 (0.8, 3.9)   |
| severe dyspepsia\(^c\)       | 0.7                | 0.9                           | 0.8 (0.3, 2.6)   | 0.4                | 0.3                    | 1.4 (0.2, 7.8)   |
| Headache                     | 1.0                | 1.5                           | 0.6 (0.3, 1.49)  | 0.7                | 0.7                    | 1.0 (0.3, 3.4)   |
| Dizziness                    | 1.5                | 2.0                           | 0.8 (0.4, 1.4)   | 0.4                | 0.9                    | 0.5 (0.1, 2.1)   |
| Tinnitus                     | <0.1               | 0.0                           | NA               | 0.3                | 0.0                    | NA               |
| Sign of overdose             | 2.6                | 3.5                           | 0.7 (0.4, 1.2)   | 1.4                | 1.7                    | 0.8 (0.3, 2.0)   |
| Other bleeding (non-GI, non cerebral) | <0.1 | 0.2 | NA | 0.1 | 0.0 | NA |
| Oral complications\(^c\)     | 0.2                | 0.8                           | 0.2 (0.1, 1.1)   | 0.1                | 0.1                    | 1.0 (0.1, 12.2)  |

\(^a\) MedDRA term.

\(^b\) Minor GI disorders include heartburn, nausea, vomiting and abdominal pain. Other terms are defined in table I.

\(^c\) Combined term.

ASA arm = ASA alone or combined with additional therapy; GI = gastrointestinal; MedDRA = Medical Dictionary for Regulatory Activities; NA = not available; OR = odds ratio; placebo arm = placebo alone or combined with the additional therapy (pseudoephedrine, lidocaine or dextromethorphan).
or anticoagulants. Patients at risk (elderly patients or those with an ulcer history) were excluded in these clinical trials.

Finally, the very low incidence of serious AEs in our study population limited our ability to investigate differences between ASA and placebo or other comparators in this regard. Much larger studies would be required to generate enough events to make such an analysis feasible.

Conclusion

In conclusion, this meta-analysis of clinical trials of short-term ASA, mostly single dose and of 1-day duration, used for the treatment of pain, fever, or colds at common OTC doses, has shown a low incidence of AEs. There was a small increase in the risk of mild to moderate dyspepsia and abdominal pain with ASA, but no significant occurrence of major GI complications was observed.

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