Gastrointestinal injury associated with NSAID use: a case study and review of risk factors and preventative strategies

Jay L Goldstein1
Byron Cryer2

1Department of Medicine, NorthShore University HealthSystem, Evanston, IL, USA; 2Division of Gastroenterology, University of Texas Southwestern Medical Center and Dallas VA Medical Center, Dallas, TX, USA

Abstract: Nonsteroidal anti-inflammatory drugs (NSAIDs) are effective anti-inflammatory and analgesic agents and are among the most commonly used classes of medications worldwide. However, their use has been associated with potentially serious dose-dependent gastrointestinal (GI) complications such as upper GI bleeding. GI complications resulting from NSAID use are among the most common drug side effects in the United States, due to the widespread use of NSAIDs. The risk of upper GI complications can occur even with short-term NSAID use, and the rate of events is linear over time with continued use. Although gastroprotective therapies are available, they are underused, and patient and physician awareness and recognition of some of the factors influencing the development of NSAID-related upper GI complications are limited. Herein, we present a case report of a patient experiencing a gastric ulcer following NSAID use and examine some of the risk factors and potential strategies for prevention of upper GI mucosal injuries and associated bleeding following NSAID use. These risk factors include advanced age, previous history of GI injury, and concurrent use of medications such as anticoagulants, aspirin, corticosteroids, and selective serotonin reuptake inhibitors. Strategies for prevention of GI injuries include anti-secretory agents, gastroprotective agents, alternative NSAID formulations, and nonpharmacologic therapies. Greater awareness of the risk factors and potential therapies for GI complications resulting from NSAID use could help improve outcomes for patients requiring NSAID treatment.

Keywords: side effects, ulcer, GI bleed, NSAID, gastrointestinal

Case study
A 53-year-old otherwise healthy female was admitted to the emergency department following two bouts of hematemesis and a single melenic stool. She denied abdominal pain or discomfort and reported no personal or family history of gastric ulcer. The patient reported being prescribed naproxen 500 mg twice daily for the 2 days prior for an ankle sprain. On examination, the patient was hypotensive in the supine position, with a blood pressure of 90/30 mmHg, and was tachycardic, with a heart rate of 130 beats per minute. Abdominal examination was benign without tenderness. Hemoglobin was 10.2 g/dL and hematocrit was 33.4%; all other evaluated laboratory values were within normal limits. Endoscopy revealed a 1×1 cm hemorrhagic gastric ulcer in the antrum with a visible vessel (Figure 1), which was cauterized at the time of endoscopy. Biopsies of the antrum and body were negative for Helicobacter pylori. Cautery was successful, and the patient was treated with an intravenous proton-pump inhibitor (PPI) and remained hospitalized for observation and to evaluate for rebleeding. During hospitalization, the patient was transitioned to an oral PPI. After 2 days
NSAID use has been associated with cardiovascular (CV), renal, and gastrointestinal (GI) complications, and certain patients are at increased risk. NSAID use results in small but consistent increases in the risk of CV events such as myocardial infarction, affected in part by dose and potency of cyclooxygenase-2 (COX-2) inhibition. NSAID use has also long been associated with kidney disease, resulting in both acute and chronic impairments in kidney function. These complications prompted the US Food and Drug Administration (FDA) to release a scientific statement in 2005 emphasizing “the importance of using the lowest effective dose for the shortest duration possible if treatment with an NSAID is warranted for an individual patient.”

More common than CV and renal complications associated with NSAID use are NSAID-related GI events, which are among the most common drug-related side effects in the United States and occur in both prescription and OTC NSAID users. These complications include bleeding gastric or duodenal ulcers and, to a lesser extent, obstructions and perforations. In a retrospective analysis of a rheumatoid arthritis patient database published in 2000, OTC ibuprofen and naproxen users had a relative risk for serious GI complications of approximately 3.5 compared with NSAID nonusers, and it is estimated that 1%–2% of continuous NSAID users experience a clinically significant upper GI event per year. These findings represent a significant clinical concern, as patients taking NSAIDs experience a relative risk of upper GI bleeding and perforations of up to 4.7 compared with nonusers.

GI complications are generally thought to be mediated primarily through inhibition of mucosal cyclooxygenase-1 (COX-1) and resultant suppression of prostaglandin production. However, COX-2 inhibition and other mechanisms, such as changes in the bacterial microbiome in the gut or the generation of free radicals, are also possibly involved. There is some question regarding the extent to which differential inhibition of COX-1 and -2 affects the GI risks associated with various NSAIDs. NSAIDs exhibit differential COX-1 and -2 inhibition and have been associated with different risks of GI and CV adverse events that vary among patients, but data sufficient to justify differences in labeling among NSAIDs in the United States have not been established.

In contrast to the acute effects of NSAIDs on the GI tract, there is some evidence that NSAID use may reduce risks of GI cancers, including gastric, pancreatic, and colorectal cancers. For example, several studies have found that non-aspirin NSAID treatments are associated with decreased risk of gastric cancer and, in the case of celecoxib, increased regression of precancerous gastric lesions compared with placebo. However, further study is needed to better characterize these potentially protective effects.
Upper GI complications

It is often noted that potentially serious GI complications commonly develop with no clinical warning symptoms suggestive of ulcers or bleeding. However, although NSAID users report increased frequency of various symptoms including reflux, belching, bloating, and/or nausea compared with nonusers, these symptoms do not reliably indicate the presence of significant upper GI mucosal injury, which includes ulcers, bleeding, perforation, obstruction, and extensive erosions. A prospective observational study found that bleeding complications occurred without typical ulcer symptoms (epigastric pain or dyspepsia) in up to 80% of affected patients. In addition, a meta-analysis of studies performed before 1997 and from 1997–2008 found that, while the overall mortality rate from GI bleeds has fallen since the mid-1990s, NSAID users with upper GI bleeding or perforation exhibit a higher mortality rate from these injuries compared with nonusers with comparable clinical scenarios. Endoscopic techniques are frequently used to manage peptic ulcer bleeding, but little research has been done to investigate whether NSAID use impacts the rate of successful hemostasis following endoscopic therapy. A retrospective study of only 76 patients found no association between NSAIDs and failure of endoscopy therapy for the treatment of gastric ulcer-associated bleeding, but the sample size was small. Interestingly, a randomized controlled trial of 224 patients who developed ulcer complications following NSAID use found that use of a COX-2 selective NSAID (celecoxib) with no effect on platelet aggregation was associated with a lower risk of recurrent bleeding compared with PPI and nonselective NSAID co-therapy; however, both therapies conferred a significant risk of rebleeding. These results led to a recommendation by the American College of Physicians that patients with previous ulcer bleeding who require an NSAID be treated with “the combination of a PPI and a COX-2 inhibitor.”

Lower GI complications

The rate of lower GI complications resulting from NSAID use has not been as widely documented as that of upper GI damage, but such complications have been recognized for decades. These injuries include bleeding in the large and small bowel, strictures of the small bowel, or exacerbation of existing illnesses such as inflammatory bowel disease. While the incidence of lower GI injury associated with NSAID use is somewhat lower than that of upper GI injury, results from a 2003 prospective study of rheumatoid arthritis patients found that 0.9% of patients taking naproxen 500 mg twice daily developed serious NSAID-associated GI injury over a 1-year period, and a 2005 retrospective study of health records in Spain found that, while the greater incidence of upper GI events results in more fatalities overall, upper and lower GI events have similar mortality rates. Video capsule endoscopy assessment of the small bowel has allowed clinicians to better quantify NSAID-related small intestinal mucosal injury, as shown by a study that found that healthy volunteers who received either naproxen 500 mg twice daily and omeprazole 20 mg once daily or celecoxib 200 mg twice daily for 2 weeks exhibited significantly more mucosal breaks compared with those receiving placebo. Slow-release diclofenac also resulted in new small intestinal mucosal injury compared with baseline after 7 days of use in almost three-quarters of healthy volunteers assessed via capsule endoscopy.

It is hypothesized that asymptomatic small bowel mucosal injury may lead to occult blood loss over time, resulting in decreases in hemoglobin levels. Results from the CONDOR (celecoxib versus omeprazole and diclofenac in patients with Osteoarthritis and Rheumatoid arthritis) study, which compared celecoxib 200 mg twice daily with diclofenac slow-release 75 mg twice daily plus omeprazole (a PPI) 20 mg once daily in arthritis patients at high risk of upper GI complications, support this concept. In that study, investigators found that, while upper GI events did not differ among treatment groups, use of diclofenac and omeprazole resulted in 3.4% of patients exhibiting a ≥2.0 g/L decrease in hemoglobin over approximately 6 months in the absence of overt bleeding. This finding suggests that GI blood loss may have been more related to slow occult bleeding. Together, these studies suggest that lower bowel injuries should be taken into account when considering the risks of NSAID use and when considering managing long-term risk.

Risks of NSAID-associated GI injuries over time

The risk of NSAID-associated GI complications is dose dependent and remains linear over time, based on the results of randomized controlled trials. The 6-month Misoprostol Ulcer Complication Outcomes Safety Assessment (MUCOSA) study, which investigated the effects of misoprostol co-administration on GI complications associated with NSAID use, and the Vioxx Gastrointestinal Outcomes Research (VIGOR) study, which compared the GI risks of naproxen with rofecoxib, both found that the cumulative incidence of upper GI events in the nonselective NSAID treatment arms was linear over time (Figure 2). In addition, two
6-month, double-blind, prospective, randomized clinical trials investigating lower GI injury following NSAID use found that the rate of patients meeting the endpoint of a decrease in hemoglobin >2 g/dL was roughly constant over time.47 Confirming these results in a clinical practice setting was the Gastrointestinal Randomized Event and Safety Open-label Non-steroidal anti-inflammatory drug Study (GI-REASONS), a randomized, prospective, open-label study comparing celecoxib with nonselective NSAIDs in osteoarthritis (OA) patients, which also found that the cumulative incidence of clinically significant upper and lower GI events was roughly linear over the 6-month trial period.15 Because GI complications associated with NSAID use are dose related and can occur at any time following exposure,32 several international societies, including the National Institute for Health and Care Excellence (NICE) in the United Kingdom, the American College of Gastroenterology, and the American College of Rheumatology (ACR), have recommended identification of risk factors and prophylaxis independent of the presence or absence of symptoms in patients with moderate-to-high risk of GI complications (Table 1).38–50

**Table 1** Characteristics of patients with an elevated risk for NSAID-associated gastrointestinal complications

<table>
<thead>
<tr>
<th>Characteristics</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Age &gt; 60 years53,54</td>
</tr>
<tr>
<td>• Helicobacter pylori infection57</td>
</tr>
<tr>
<td>• Receiving hemodialysis52</td>
</tr>
<tr>
<td>• High-dose or multiple NSAID use55</td>
</tr>
<tr>
<td>• History of upper gastrointestinal injury14,51</td>
</tr>
<tr>
<td>• Anticoagulant use56,59</td>
</tr>
<tr>
<td>• Oral corticosteroid use; prednisone55,60</td>
</tr>
<tr>
<td>• Selective serotonin reuptake inhibitor use61,62,64</td>
</tr>
</tbody>
</table>

**Abbreviation:** NSAID, nonsteroidal anti-inflammatory drug.

**Risk factors for NSAID-associated GI injury**

A variety of patient characteristics are associated with increased risk for NSAID-related GI complications (Table 1). Patients with a history of GI injury are at higher risk for GI complications following NSAID use,14,51 and patients with renal failure who are on hemodialysis also exhibit increased risk of GI bleeding with NSAID use.52 Age is an important factor, with risk increasing with increasing age. As the absolute risk varies by age, a threshold of risk based on age is often suggested to be >60 years old.53,54 Patients taking high-dose NSAIDs and those taking NSAIDs with aspirin, even at low, CV-prevention doses (≤325 mg/day), have elevated risks of GI events.55 Certain medications also increase the risk of GI injury when used concurrently with NSAIDs. For example, use of oral corticosteroids coadministered with NSAIDs is associated with an increase in the rate of GI complications as much as twofold compared with patients taking NSAIDs alone.55 Anticoagulants have been found to substantially increase the risk for elderly patients of developing ulcer...
bleeding when used with NSAIDs. Additionally, a Danish study of prescription and hospitalization records of patients ages 16 to 105 years found that anticoagulants and non-salicylate NSAIDs taken concurrently increased upper GI bleeding more than anticoagulants taken with aspirin or acetaminophen. Furthermore, the increased risk of ulcer bleeding due to anticoagulant use may not be mitigated by gastroprotective agents. Selective serotonin reuptake inhibitors also increase risk of upper GI complications when used with NSAIDs, as several studies have shown that concurrent selective serotonin reuptake inhibitor and NSAID use results in a greater increase in the incidence of GI bleeding than the sum of their independent effects would suggest. These results suggest that caution should be used when considering prescribing NSAIDs to patients using these agents.

The limited awareness of risk factors results in many patients receiving inadequate preventative therapies. For example, a study of veterans prescribed NSAIDs over a 1-year period showed that nearly three-quarters (73%) of the patients with risk factors for NSAID-related upper GI injury were not receiving appropriate gastroprotective therapy based on evidence-based guidelines. In fact, prescription practices may frequently be inappropriate when a patient’s GI and CV history are considered, according to results from a Spanish National Health System study conducted in 2011, which found that 74% of OA patients with elevated risk for GI and CV NSAID-related side effects were receiving nonselective NSAIDs or COX-2 selective NSAIDs. These data indicate that not only do NSAIDs represent heightened risks to some patients, but that awareness of the risk factors and of the use of preventative therapy for NSAID-related upper GI injury could be improved.

**Approaches to the prevention of GI injuries from NSAIDs**

PPIs and histamine-2 (H2) receptor antagonists

Coadministration of NSAIDs with PPIs is a well-documented and effective, although underutilized, approach to reduce endoscopic damage and control dyspeptic symptoms associated with the use of NSAIDs (Table 2). Infrequent side effects associated with PPIs have occurred; these may include an increased chance of pneumonia compared with nonusers, hypomagnesemia, increased incidence of spine and hip fractures, as well as an increased chance of contracting *Clostridium difficile*-associated diarrhea compared with PPI-naïve patients. H2 receptor antagonists (H2RAs), which inhibit acid secretion, have also been evaluated for reducing NSAID-associated complications. A meta-analysis of 14 trials found that H2RAs (eg, famotidine and ranitidine) were protective at high doses, but, at commonly prescribed doses, they reduced the risk of duodenal but not gastric ulcers. This was demonstrated in a 24-week study of patients requiring chronic NSAID treatment, which showed that coadministration of high-dose (80 mg/day) famotidine with ibuprofen (2,400 mg/day) resulted in significantly fewer gastroduodenal ulcers compared with ibuprofen alone. Rare side effects associated with H2RA use include increased risk of pneumonia.

**Misoprostol**

Co-prescribed misoprostol, a synthetic prostaglandin used to counteract the prostaglandin-inhibitory actions of NSAIDs, decreases NSAID-related upper GI complications by approximately 40% compared with NSAIDs alone. The single-tablet combination therapy of diclofenac sodium–misoprostol

Table 2 Gastroprotective strategies to prevent gastrointestinal complications associated with NSAID use

<table>
<thead>
<tr>
<th>Preventative strategy</th>
<th>Gastroprotection strategy</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>Proton-pump inhibitors</td>
<td>Reduce secretion of gastric acid</td>
<td>Associated with increased risk of pneumonia, hypomagnesemia, <em>Clostridium difficile</em>-associated diarrhea, and spine and hip fractures</td>
</tr>
<tr>
<td>Misoprostol</td>
<td>Synthetic prostaglandin that counteracts prostaglandin inhibition by NSAIDs</td>
<td>Gastrointestinal discomfort, nausea, and diarrhea; fetal loss</td>
</tr>
<tr>
<td>Histamine-2 receptor agonists</td>
<td>Reduce secretion of gastric acids</td>
<td>Increased risk of pneumonia</td>
</tr>
<tr>
<td>COX-2 inhibitors</td>
<td>Selective for COX-2 receptor, which circumvents the inhibition of gastroprotective prostaglandins by COX-1</td>
<td>Cardiovascular complications similar to those associated with other NSAIDs</td>
</tr>
<tr>
<td>Topical formulations</td>
<td>Lowers systemic drug exposure to potentially reduce risk</td>
<td>Impractical for systemic pain and may be ineffective for long-term pain relief</td>
</tr>
<tr>
<td>Lower-dose oral formulations</td>
<td>Lowers systemic drug exposure to reduce risk</td>
<td>Limited efficacy and safety data</td>
</tr>
</tbody>
</table>

**Note:** Only effective as a protective agent at 300 mg twice daily for ranitidine, ≥40 mg twice daily for nizatidine, and 80 mg total daily dose for famotidine.

**Abbreviations:** COX, cyclooxygenase; NSAID, nonsteroidal anti-inflammatory drug.
was found to be an effective gastroprotective therapy measured by cost per life-year gained in a study in the Netherlands.\textsuperscript{84} Of note is that the single-tablet formulation may improve patient adherence, as nonadherence with multiple-pill therapeutic regimens increases the risk of GI events.\textsuperscript{69,85,86} Unfortunately, misoprostol is not always well tolerated due to diarrhea and abdominal pain, preventing continued use; lower doses of misoprostol with a lesser frequency of side effects may also be less effective at preventing GI events.\textsuperscript{74} Misoprostol is an abortifacient, contraindicating its use in women who are pregnant or may become pregnant (Table 2).\textsuperscript{75} Rebamipide is a cytoprotective anti-ulcer drug that stimulates prostaglandin production. At a dose of 100 mg three times daily (TID), it was found to be significantly more effective in reducing rates of endoscopic gastric or duodenal ulcer compared with misoprostol 200 mg TID in the Study of NSAID-induced GI Toxicity Prevention by Rebamipide and Misoprostol (STORM), a multicenter, 12-week, randomized controlled trial of patients using NSAIDs. Rebamipide is not approved for use in the United States.\textsuperscript{87}

**COX-2 inhibitors**

COX-2 selective inhibitors are associated with less risk of GI injury than nonselective NSAIDs due to lesser inhibition of COX-1, which is involved in the maintenance of gastric mucosa integrity. COX-2 selective inhibitor use was associated with a decrease in the risk of symptomatic ulcers and clinically significant ulcer complications compared with nonselective NSAIDs, according to a 2007 meta-analysis.\textsuperscript{88} In the CONDOR study, while efficacy was similar, fewer patients treated with the COX-2 selective inhibitor celecoxib experienced reductions in hemoglobin or withdrew from the study due to adverse GI events compared with those treated with diclofenac plus omeprazole.\textsuperscript{44,45} However, use of COX-2 inhibitors at therapeutic doses may increase CV risks.\textsuperscript{77} Following an FDA warning about the CV risks of COX-2 inhibitors and all nonselective NSAIDs, and the voluntary withdrawals of rofecoxib and valdecoxib from the market,\textsuperscript{49} prescriptions for COX-2 inhibitors decreased by 12% and those for nonselective NSAIDs increased by 47% between 2003 and 2005. However, this trend was not accompanied by an increase in prescriptions of gastroprotective co-therapies.\textsuperscript{90}

**Enteric-coated NSAIDs**

While at least one study found that enteric-coated NSAIDs reduce upper GI events,\textsuperscript{91} most data indicate that enteric-coated NSAIDs do not reduce incidence of upper GI events compared with other formulations.\textsuperscript{92–94} Interestingly, attempts to reduce upper GI symptoms through use of slow-release and enteric-coated formulations may hypothetically increase lower GI complications.\textsuperscript{95}

**Topical NSAIDs**

Because NSAID-associated GI complications are dose dependent, development of formulations that lower systemic exposure while providing efficacious pain relief may reduce GI injury. Topical NSAID formulations can produce higher concentrations of drug in local tissue with very low systemic exposure as measured via plasma concentrations,\textsuperscript{96} and use of topical NSAIDs may be associated with fewer GI events (Table 2).\textsuperscript{97–101} Although topical NSAID formulations have been shown to be effective in treatment of acute pain,\textsuperscript{79} and for short-term use in treating chronic pain, there are conflicting results regarding whether topical NSAIDs provide effective long-term pain relief.\textsuperscript{102,103} Further study is necessary to determine the long-term benefits and risks of topical NSAID use.

**Lower-dose NSAID formulations**

New formulations of NSAIDs may reduce risks of adverse events by using lower doses while providing effective analgesia (Table 2). There is some evidence that some NSAIDs, such as diclofenac, could provide effective pain relief at lower doses than are currently used, assuming 80% inhibition of COX-2 is necessary for therapeutic efficacy.\textsuperscript{51} This would hypothetically provide effective pain relief with an improved GI safety profile due to lessened inhibition of COX-1.\textsuperscript{29} A diclofenac potassium liquid-filled capsule\textsuperscript{64} using a formulation designed to deliver diclofenac more rapidly than conventional tablets was approved by the FDA in 2009.\textsuperscript{105,106} Absorption of the liquid-filled capsules is faster than that of diclofenac potassium immediate-release tablets, and the capsules produce greater pain relief compared with placebo at lower doses of diclofenac (25 mg four times daily) than are generally used;\textsuperscript{79,105,107} however, it is unclear whether they produce more rapid or effective pain relief than other diclofenac formulations. Lower-dose capsules that contain finely milled, rapidly absorbed NSAID particles may also provide analgesia at lower systemic doses.\textsuperscript{90,81} Low-dose diclofenac capsules (18 mg or 35 mg TID for mild-to-moderate pain) and indomethacin capsules (20 mg TID or 40 mg two times daily or TID) containing fine-milled particles have been approved by the FDA for treatment of mild-to-moderate acute pain in adults\textsuperscript{25,108} and have been found to provide effective relief of acute, postoperative pain in Phase III studies.\textsuperscript{80,81} Additionally, low-dose diclofenac (35 mg TID) has been shown to
provide effective treatment of OA pain in a 12-week study and has been approved by the FDA for management of OA-related pain. A low-dose naproxen capsule was also found to effectively relieve postoperative dental pain in a Phase II study.

**H. pylori**

Eradicating *H. pylori* may decrease GI risks in some NSAID users, which could reduce worldwide incidence of NSAID-related GI injury, as *H. pylori* affects up to 50% of the worldwide population. One systematic literature review found that *H. pylori* eradication in infected patients was as effective as the use of PPIs in preventing GI complications due to NSAID use; however, another found that, although *H. pylori* eradication reduces risk, PPIs provided superior ulcer prevention. While it is unclear whether *H. pylori* eradication is as effective as other strategies, it may provide benefit to some patients.

**Nonpharmacologic therapies**

Another possibility for reducing the incidence of NSAID-associated GI complications is to reduce NSAID use through adoption of alternative therapies. While assessment of their effectiveness is challenging, therapies such as acupuncture and physical therapy/exercise may provide relief for some patients. While some randomized controlled trials have found acupuncture to be more effective for OA pain relief than sham treatments, a meta-analysis of eleven studies published between 1994 and 2006 found sufficiently heterogeneous results that the authors were unable to draw firm conclusions regarding acupuncture’s efficacy. While significant results have been found for use of acupuncture, particularly for knee OA, the effect is generally small, and larger studies are needed. Exercise and physical therapy may also provide pain relief, as they have been found to improve pain and function in knee OA, but may delay the need for surgical intervention and may reduce the need for medication. Because of these results, the ACR has issued guidelines strongly recommending exercise for knee OA. Unfortunately, the effect of exercise on knee OA may be short-term, and the extent of functional improvement is unclear. While many approaches for prevention of NSAID-associated GI injury show effectiveness in some studies, practical considerations prevent their universal use.

**Cost-effectiveness**

The direct cost of preventative strategies to patients and payers and the absolute patient risk for GI complications are the key factors that influence cost-effectiveness. The relative cost of preventing a single complication is high in low-risk populations and is the basis of recommendations from the ACR and other health authorities that indicate that low-risk patients should not receive gastroprotective therapies. The picture becomes more complicated in patients with higher risk for GI injury; as risk increases, the associated costs of prescribing such therapies is increasingly offset by the escalating cumulative costs associated with the adverse events. Net costs/savings are driven by the additional procurement costs of the preventative therapy and the expected frequency of adverse events based on risk. Many studies have tried to evaluate cost-effectiveness, but it is difficult to make decisions based on finances given that the cost of PPIs has dramatically declined and that PPIs are now available OTC. Additionally, cost-effectiveness studies do not always adequately take into account the impact of injury on quality of life. An economic model examining PPI use in three large randomized trials, weighted by quality of life, found that use of PPIs with either COX-2 selective inhibitors or nonselective NSAIDs was cost-effective in OA patients, even those at low risk of GI events, with the caveat that the mean age of participants was above 60 years and thus these patients may not be considered to be objectively low risk. These studies suggest that the economic picture of how to most cost-effectively decrease NSAID-associated GI injury is not yet clear and that, together, studies considering different risk pools are necessary to determine optimal management of patient subpopulations.

**Balancing risks**

The CV risks associated with NSAID use have received increased attention recently, such that the American Heart Association released a statement declaring that NSAID use should be “coupled with the realization that effective pain relief may come at the cost of a small but real increase in risk for cardiovascular or cerebrovascular complications.” In February 2014, the FDA reviewed the data surrounding the CV complications associated with NSAID use and determined that there were insufficient data to distinguish CV toxicity among individual NSAIDs, including COX-2 selective inhibitors and naproxen, and that this class of agents was associated with an increased risk of ischemic events. This attention to the risks associated with NSAIDs and the potential differences among specific NSAIDs represents a growing awareness about the complications associated with this class of drugs.
Conclusion
GI mucosal injury associated with use of NSAIDs is a serious clinical concern, and studies suggest that the rate of complications does not decrease with duration of use. There are several strategies and NSAID drug product formulations that may be associated with decreased GI risk, but there is no one therapy that will provide optimal pain relief and decrease risk for all patients. Also, although nonpharmacological therapies have promise, often they have been inadequately studied compared with pharmacological therapies. Meanwhile, the high cost of GI events to the health care system and to patients’ quality of life necessitates improvement in the risk–benefit profile of NSAIDs or development of alternative medications or therapies. In addition, the CV and renal side effects of NSAIDs must be considered alongside reducing the risk of GI complications. Optimally, developments in pain management will focus on tailoring therapies to the individual patient. Also, in addition to development of new therapies, improvements in patient and provider education and patient adherence are necessary to improve outcomes. Greater awareness of the short-term GI risks of NSAIDs, including potential overuse of OTC NSAIDs and more frequent use of gastroprotection, might have prevented the ulcer in the patient described in the case study at the beginning of this article.

Acknowledgments
Editorial support for this manuscript was provided by Jill See, PhD, and Colville Brown, MD, of AlphaBioCom LLC (King of Prussia, PA, USA). Funding for editorial support was provided by Iroko Pharmaceuticals, LLC (Philadelphia, PA, USA).

Disclosure
JLG has served as an advisory board attendee for Iroko Pharmaceuticals, LLC. BC has served as a consultant for Iroko Pharmaceuticals, LLC; Ritter Pharmaceuticals; Sanofi Pharmaceuticals; Sandoz Pharmaceuticals; and Sucampo, Inc. The authors report no other conflicts of interest in this work.

References


