Efficacy and tolerability of lumiracoxib, a highly selective cyclo-oxygenase-2 (COX2) inhibitor, in the management of pain and osteoarthritis

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Abstract: Lumiracoxib is a COX2 inhibitor that is highly selective, is more effective than placebo on pain in osteoarthritis (OA), with similar analgesic and anti-inflammatory effects as non-selective NSAIDs and the selective COX2 inhibitor celecoxib, has a lower incidence of upper gastrointestinal (GI) side effects in patients not taking aspirin, and a similar incidence of cardiovascular (CV) side effects compared to naproxen or ibuprofen. In the context of earlier guidelines and taking into account the GI and CV safety results of the TARGET study, lumiracoxib had secured European Medicines Agency (EMEA) approval with as indication symptomatic treatment of OA as well as short-term management of acute pain associated with primary dysmenorrhea and following orthopedic or dental surgery. In the complex clinical context of efficiency and safety of selective and non-selective COX inhibitors, its prescription and use should be based on the risk and safety profile of the patient. In addition, there is further need for long-term GI and CV safety studies and general post-marketing safety on its use in daily practice. Meanwhile, at the time of submission of this manuscript, the EMEA has withdrawn lumiracoxib throughout Europe because of the risk of serious side effects affecting the liver.

Keywords: lumiracoxib, NSAIDs, COX2 inhibitors, gastro-intestinal and cardiovascular safety

Introduction
Non-selective non-steroidal anti-inflammatory drugs (NSAIDs) are widely used by patients with acute pain or with chronic pain due to osteoarthritis (OA) and rheumatoid arthritis (RA) (ACR Recommendations 2000; Hochberg 2002; Jordan et al 2003; Schnitzer 2006). They put them at increased risk for clinically important damage to the mucosa of the upper part of the gastrointestinal tract (Wolfe et al 1999), with a relative risk (RR) of 2.7–5.4 for the so-called PUBs and POBs (PUBs: Perforation, clinically manifest Ulcer and Bleeding; POBs: Perforation, Obstruction, Bleeding) (Hernandez-Diaz and Rodriguez 2000; Ofman et al 2002). OA and RA patients were found to be 2.5–5.5 times more hospitalized for NSAID-related gastro-intestinal (GI) events than the general population (Singh 1988) and 5%–10% of PUBs are fatal (Armstrong and Blower 1987).

Cyclo-oxygenase 2 (COX2)-selective inhibitors have been developed based on the finding that the COXs that are constitutively involved in the physiology of the GI mucosa (COX1) are different from the ones that are inducible by inflammation (COX2) (Vane et al 1994; Warner et al 1999; Fitzgerald 2003). Therefore, selective inhibition of COX2 could dissociate anti-inflammatory activity from GI side effects. In this way, COX2 selective inhibitors should reduce clinically manifest ulcers and ulcer complications compared with non-selective NSAIDs, meanwhile exerting similar effects on acute pain.
vascular thrombotic events (Solomon 2005). The use of COX2 inhibitors induces an elevated risk of thrombo-embolic events (Fitzgerald 2003). It has been suggested that COX2 is upregulated in vascular segments for thrombotic events, including stroke and particularly myocardial infarction (MI), as demonstrated in the VIGOR study, in which the risk of MI was higher in RA-patients treated with rofecoxib than in naproxen users (Bombardier et al 2000). It was debated whether this difference in CV risk was the result of a protective effect of naproxen on the incidence of MI (Bombardier et al 2000), or a consequence (side-effect) of the use of COX2 inhibitors.

Later on, in studies developed to observe whether the use of COX2 inhibitors protected against the occurrence of colonic polyps, it has become clear that rofecoxib, as compared to placebo, doubles the risk for thrombotic events, mainly myocardial infarction and ischemic CV events (Bresalier et al 2005; Kerr et al 2007). Also for other COX2 inhibitors, celecoxib and valdecoxib, an elevated CV risk has been shown (Nussmeier et al 2005; Solomon et al 2005).

Although the exact mechanism of this association is still unclear, the balance between prostacyclin (PGI2) and thromboxane A2 (TXA2) is presumably shifted to an increased risk for thrombo-embolic events (Fitzgerald 2003). It has been suggested that COX2 is upregulated in vascular segments under conditions of increased vascular shear stress, and that the reduction of endovascular production of prostacyclin by the use of COX2 inhibitors induces an elevated risk of vascular thrombotic events (Solomon 2005).

Besides an elevated CV risk, it has recently been shown that the use of conventional NSAIDs is associated with edema, congestive heart failure, cardiac arrhythmias, and an increased risk of CV events (Kearney et al 2006; McGettigan and Henry 2006; Vonkemann et al 2006; Zhang et al 2006). In this context, it was realised that the development of new selective COX2 inhibitors had to be evaluated in a much broader context of safety than only GI protection and had to include data on CV safety (Silverstein et al 2000; Farkouh et al 2004; Schnitzer et al 2004; Cannon et al 2006; Laine et al 2007). In order to perform such studies, larger trials were needed with more endpoints than in the original studies that focused on GI protection only. In addition, also for the non-selective NSAIDs additional data were required to adequately judge their effect on CV risk.

Several clinical trials have been performed to study both the GI and CV safety with COX2 (Tables 1–4), including VIGOR (rofecoxib vs naproxen) (Bombardier et al 2000), CLASS (celecoxib vs diclofenac and ibuprofen) (Silverstein et al 2000), MEDAL (etoricoxib vs diclofenac in a pooled analysis) (Cannon et al 2006; Laine et al 2007), and TARGET (lumiracoxib vs ibuprofen and naproxen) (Farkouh et al 2004; Schnitzer et al 2004), which will be discussed in detail later in this review.

These studies differed between each other in many aspects. Differences included indications for treatment (OA and/or RA), number of patients included (between 8059 and 37,701), the comparator non-selective NSAIDs (naproxen, diclofenac, ibuprofen; none was placebo-controlled), duration of the study (6–36 months), exclusion or not because of history of GI risks/events (GI surgery, current bleeding, active GI disorder, recent ulcer, intake of GI protectors, bleeding last year, any perforation, obstruction or not), GI protection during the study (not allowed, allowed, stimulated or not), GI protection (liberal for placebo), definitions of GI end-points (various combinations of PUBOs, eventually further specified as complicated or not), exclusion or not because of history of CV risks/events (during variable time periods before the study (during last 6 months to ever), and defined as any CV event or specified as MI (clinical or on ECG), coronary bypass, percutaneous coronary intervention, stroke, new angina, low dose aspirin (allowed, stimulated, or not allowed), severe heart failure), and definitions of CV side effect endpoints (MI alone, composite endpoints including MI, stroke, angina, any or arterial thrombotic events, including more standardized outcomes from the Anti-Platelet Trialists’ Collaboration endpoint).
In these studies, rofecoxib and etoricoxib decreased the risk of GI event endpoints (Table 2), but the size and significance of effects according to definitions of GI events and reporting on significance of interaction of treatment-by-subgroup analysis differed between the drugs (Bombardier et al 2000; Silverstein et al 2000; Cannon et al 2006; Laine et al 2007). Celecoxib did not decrease the primary GI endpoint (POBs) over the entire duration of the study, but significantly reduced the risk of GI events when symptomatic ulcers were included in the analysis (Silverstein et al 2000).

In these studies, rofecoxib increased the risk of MI (HR: 4.25, CI: 1.39–17.37) (Bombardier et al 2000), which was the reason for withdrawal of rofecoxib (Vioxx®) from the market in September 2004. Low dose celecoxib and etoricoxib did not affect the risk of pre-specified composite CV endpoints in the total group of patients and in specified subgroups (on low dose aspirin or not) as compared to their comparator non-selective NSAID(s) (Table 3) (Silverstein et al 2000; Cannon et al 2006; Laine et al 2007).

It is important to realize that the similarity of incidence of CV risk between selective and non-selective COX inhibitors does not mean that there is no increased CV risk of selective COX2 inhibitors compared to placebo. On the contrary, compared to placebo, it has been demonstrated that the selective COX2 inhibitors rofecoxib (Bresalier et al 2005; Kerr et al 2007) and celecoxib (Solomon et al 2005) at high doses increase the risk of CV events compared to placebo. Furthermore, non-selective NSAIDs, such as ibuprofen (Antman et al 2007), diclofenac (Antman et al 2007), but probably not naproxen (Antman et al 2007) and even paracetamol (Chan et al 2006), were associated with an increased risk for CV events compared to placebo in meta-analyses.

In this complex safety context, we review here the data on the effects and safety of lumiracoxib, a highly selective COX2 inhibitor.

**Lumiracoxib**

**Preclinical data**

Lumiracoxib (Prexige®) is a selective COX2 inhibitor developed for the treatment of osteoarthritis, rheumatoid arthritis, and acute pain (Lyseng-Williamson and Curran 2004; Esser et al 2005). Lumiracoxib differs structurally from other drugs in the class of selective COX2 inhibitors (Brune and Hinz 2004; Mangold et al 2004). The other inhibitors contain a tricyclic ring and a sulfone or sulfonamide group (Brune and Hinz 2004) whereas lumiracoxib is a phenyl acetic acid derivative. It has the highest selectivity (selective for COX2 compared with COX1 in the human whole blood assay with a ratio of 515:1 in healthy subjects and in patients with osteoarthritis or rheumatoid arthritis) and a fairly short plasma half-life (3–6 hours) compared with other COX2-selective inhibitors (Esser et al 2005).

Lumiracoxib has good oral bioavailability (74%). It is rapidly absorbed, reaching maximum plasma concentrations 2 hours after dosing, and is highly plasma protein bound. Lumiracoxib has a short elimination half-life from plasma (mean 4 hours) and demonstrates dose-proportional plasma

<table>
<thead>
<tr>
<th>Drug (reference)</th>
<th>Study</th>
<th>Diagnoses and patients (n)</th>
<th>Duration (months)</th>
<th>Comparator drug</th>
<th>Risk patients excluded</th>
<th>CV history</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rofecoxib (Bombardier 2000)</td>
<td>VIGOR</td>
<td>RA (8 076)</td>
<td>12</td>
<td>Naproxen</td>
<td>GI surgery, current B, IBD, previous or current PPI</td>
<td>History of CV last 2 yrs, MI or coronary bypass last yr, previous or current aspirin</td>
</tr>
<tr>
<td>Celecoxib (Silverstein 2000)</td>
<td>CLASS</td>
<td>OA, RA (8 059)</td>
<td>6</td>
<td>Ibuprofen, diclofenac, Diclofenac</td>
<td>Active GI, U last 30 d, any GI surgery</td>
<td>No exclusion criteria</td>
</tr>
<tr>
<td>Etoricoxib (Cannon 2006; Laine 2007)</td>
<td>MEDAL</td>
<td>OA, RA (37 701)</td>
<td>36</td>
<td></td>
<td>No exclusion if at GI risk</td>
<td>MI or coronary bypass or percutaneous coronary intervention during last 6 mo, aspirin recommended if at CV risk</td>
</tr>
<tr>
<td>Lumiracoxib (Schnitzer 2004; Farkouch 2004)</td>
<td>TARGET</td>
<td>OA (18 325)</td>
<td>12</td>
<td>Ibuprofen, naproxen</td>
<td>On GI protection, U last 3 mo, B last yr, any P or O</td>
<td>MI or coronary bypass or percutaneous coronary intervention during last 6 mo, aspirin recommended if at CV risk, MI (clinical, on ECG), stroke, coronary bypass graft surgery, new angina during last 6 mo, high CV risk without aspirin, severe heart failure, on anticoagulation therapy</td>
</tr>
</tbody>
</table>

Abbreviations: B, bleeding; CV, cardiovascular; GI, gastrointestinal; IBD, inflammatory bowel disease; MI, myocardial infarction; O, obstruction; OA, osteoarthritis; P, perforation; RA, rheumatoid arthritis; PPI, proton pump inhibitors; U, ulcer.
pharmacokinetics with no accumulation during multiple dosing.

Lumiracoxib is metabolized extensively prior to excretion, with only a small amount excreted unchanged in urine or feces. Lumiracoxib and its metabolites are excreted via renal and fecal routes in approximately equal amounts (Lyseng-Williamson and Curran 2004).

Lumiracoxib does not exhibit any clinically meaningful interactions with a range of commonly used medications including aspirin (acetylsalicylic acid), fluconazole, an ethinylestradiol- and levonorgestrel-containing oral contraceptive, omeprazole, the antacid Maalox®, methotrexate, and warfarin (although, as in common practice, routine monitoring of coagulation is recommended when lumiracoxib is co-administered with warfarin) (Lyseng-Williamson and Curran 2004).

**Clinical efficacy**

The effectiveness of lumiracoxib was superior to placebo in patients with OA at doses of 100 mg and 200 mg once daily and similar to celecoxib and non-selective NSAID, as described and reviewed in detail in Lyseng-Williamson and Curran (2004).

Several studies have been shown the superiority of lumiracoxib compared to placebo in pain in RA (Guesens et al 2004), after surgery, orthopedic surgery, primary dysmenorrhea and tension headache (Lyseng-Williamson and Curran 2004).

**Safety**

**Upper gastrointestinal safety: endoscopic studies**

In endoscopic studies, lumiracoxib has been associated with a rate of acute gastric injury and chronic ulcer formation that does not differ from placebo (Rordorf et al 2003b) and which was significantly lower than with the non-selective NSAID ibuprofen and with celecoxib (Hawkey et al 2004; Kivitz et al 2004).

**Clinical gastrointestinal safety**

To establish the gastrointestinal safety of lumiracoxib, the Therapeutic Arthritis Research and Gastrointestinal Event Trial (TARGET) was performed to test the hypothesis that patients with osteoarthritis, randomized to lumiracoxib (400 mg once daily, which is 2–4 times the recommended dose for osteoarthritis), had significantly fewer complicated ulcers than patients randomized to either ibuprofen.

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**Table 2 Gastrointestinal endpoints in studies of selective COX2 inhibitors**

<table>
<thead>
<tr>
<th>Drug (reference)</th>
<th>Study</th>
<th>GI protection allowed in study</th>
<th>GI endpoints</th>
<th>Hazard ratios, relative risks (RR) or incidence (%) of upper GI risk endpoints</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>All</td>
<td>No aspirin</td>
</tr>
<tr>
<td>Rofecoxib (Bombardier 2000)</td>
<td>VIGOR</td>
<td>Yes</td>
<td>Confirmed POBU</td>
<td>0.5 (0.3–0.6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Complicated GI (POB)</td>
<td>0.4 (0.2–0.8)</td>
</tr>
<tr>
<td></td>
<td>CLASS</td>
<td>No</td>
<td>POB</td>
<td>RR: 0.53 (0.26–1.11)</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Symptomatic U + POB</td>
<td>RR: 0.59 (0.38–0.94)</td>
</tr>
<tr>
<td>Celecoxib (Silverstein 2000)</td>
<td></td>
<td></td>
<td>Clinical POBU</td>
<td>0.69 (0.57–0.83)</td>
</tr>
<tr>
<td></td>
<td>MEDAL</td>
<td>Yes</td>
<td>All</td>
<td>0.62 (0.45–0.83)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>No PPI</td>
<td>0.74 (0.58–0.95)</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>+ PPI</td>
<td>0.91 (0.67–1.24)</td>
</tr>
<tr>
<td>Etoricoxib (Cannon 2006; Laine 2007)</td>
<td></td>
<td></td>
<td>All</td>
<td>1.03 (0.70–1.52)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>No PPI</td>
<td>0.72 (0.42–1.22)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>+ PPI</td>
<td>0.57 (0.45–0.74)</td>
</tr>
<tr>
<td>Lumiracoxib (Schnitzer 2004; Farkouh 2004)</td>
<td>TARGET</td>
<td>No</td>
<td>Definite or probable complicated U (POB)</td>
<td>vs ibuprofen + naproxen</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>vs ibuprofen</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td>vs naproxen</td>
</tr>
</tbody>
</table>

aNo treatment-by-subgroup interaction.

**Abbreviations:** B, bleeding; GI, gastrointestinal; O, obstruction; P, perforation; PPI, proton pump inhibitors; U, ulcer.
Lumiracoxib in the management of pain and osteoarthritis

(3 × 800 mg/day) or naproxen (2 × 500 mg/day), without affecting CV risk (Farkouh et al 2004; Schnitzer et al 2004).

Table 3 Cardiovascular endpoints in studies of selective COX2 inhibitors

<table>
<thead>
<tr>
<th>Drug (reference)</th>
<th>Study</th>
<th>Aspirin allowed</th>
<th>CV endpoints</th>
<th>Hazard ratios or incidence of CV endpoints</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>All</td>
<td>No aspirin</td>
</tr>
</tbody>
</table>

Rofecoxib (Bombardier 2000)  
CLASS Yes MI vs ibuprofen + naproxen 1.14 (0.78–1.66) 1.22 (0.74–2.02) 1.04 (0.59–1.84)  
VIGOR No MI death vs ibuprofen 0.95 (0.81–1.11) 1.0% vs 1.0%, NS 1.06% vs 1.78% vs 1.87%, NS  
Celexcoxib (Silverstein 2000)  
CLASS Yes MI, stroke or angina 0.96 (0.79–1.16) 0.96 (0.79–1.16) 0.96 (0.79–1.16)  
Etoricoxib (Cannon 2006; Laine 2007)  
MEDAL Stimulated Any thrombotic events 0.96 (0.81–1.13) 1.0% vs 1.0%, NS 1.06% vs 1.78% vs 1.87%, NS  
TARGET Yes APTC (all MI, stroke or vascular death) vs ibuprofen + naproxen 1.14 (0.78–1.66) 1.22 (0.74–2.02) 1.04 (0.59–1.84)  
Lumiracoxib (Schnitzer 2004; Farkouh 2004)  
TARGET Yes APTC (all MI, stroke or vascular death) vs ibuprofen 1.14 (0.78–1.66) 1.22 (0.74–2.02) 1.04 (0.59–1.84)  
confirmed or probable MI (clinical and silent) vs naproxen 1.46 (0.89–2.37) 1.49 (0.76–2.92) 1.42 (0.70–2.90)  
vs ibuprofen + naproxen 1.31 (0.70–2.45) 1.42 (0.63–3.39) 1.14 (0.44–2.95)  
vs ibuprofen 0.66 (0.21–2.09) 0.75 (0.20–2.79) 0.47 (0.04–5.14)  
vs naproxen 1.77 (0.82–3.84) 2.37 (0.74–7.55) 1.36 (0.47–3.93)  

<sup>a</sup>Pen-protocol analysis, similar results after ITT analysis.  
<sup>b</sup>APTC, Anti-Platelet Trialists’ Collaboration endpoint.  
<sup>c</sup>Incidence.  
<sup>d</sup>Events per 100 patient yrs.  
<sup>e</sup>Abbreviation: nr, not reported.

Clinical cardiovascular safety

Based on CV risk profile at baseline, patients were excluded from the TARGET study if they had a history of MI (clinical or silent as shown on ECG), stroke, coronary bypass, new angina of recent onset (last 6 months), high CV risk without intake of aspirin or severe heart failure (Farkouh et al 2004). The primary composite endpoint (the incidence of MI, stroke, and CV death) did not differ between lumiracoxib and ibuprofen or naproxen combined (RR: 1.14, CI: 0.78–1.66) or separately, irrespective of aspirin use (RR: 1.22 and 1.04). Thus, the overall CV signal showed no difference but this is because the study is underpowered to demonstrate statistically significant and clinically meaningful differences. There was a trend towards more CV events relative to naproxen (RR: 1.46, CI: 0.89–2.37) and fewer events relative to ibuprofen (RR: 0.76, CI: 0.41–1.40).

It is important to realize that the TARGET trial has been performed in patients with a relatively low risk for CV events. Patients were excluded from the study if they had a MI, stroke, coronary artery bypass graft surgery, percutaneous coronary intervention or new-onset angina within 6 months prior to screening, electrocardiogram (ECG) evidence of silent myocardial ischemia, New York Heart Association congestive heart failure class III–IV, or if they were receiving anticoagulation therapy. As a consequence, it cannot be fully excluded that a difference between the comparators
would have been observed if more high risk patients would have been enrolled.

The incidence of composite GI and CV endpoints combined fell by 35% in patients treated with lumiracoxib compared to both comparators (RR: 0.65, CI: 0.49–0.84), by 50% compared to ibuprofen (p < 0.01) and by 25% compared to naproxen (NS).

Other safety data
Non-selective NSAIDs interfere with COX1 and COX2 in the kidney (Harris 2006). Their most common renal side effect is peripheral edema due to increased sodium retention (Harris 2006), which also is one of the contributors to an increase in blood pressure. Acute renal failure is a rare but potentially serious complication of non-selective NSAIDs (Harris 2006), which predominantly occurs in elderly with cardiovascular involvement during periods of dehydration (diarrhea, high fever). Also selective COX2 inhibitors may cause edema, congestive heart failure and modest elevations of blood pressure (Harris 2006). In the MEDAL study, more discontinuations were observed in etoricoxib users due to hypertension and edema than in diclofenac users (no difference was found for congestive heart failure) (Cannon et al 2006; Laine et al 2007).

In the TARGET study (Farkouh et al 2004; Schnitzer et al 2004), the incidence of major renal events and chronic heart failure were similar between the treatment groups (Table 4), while unfortunately the incidence of peripheral edema was not reported. Interestingly, blood pressure remained stable in patients treated with lumiracoxib, which was significantly different from the moderate but significant increase in blood pressure with naproxen and diclofenac. In the TARGET study no data were presented on cutaneous side effects (Farkouh et al 2004; Schnitzer et al 2004).

Hepatotoxicity has been the reason for withdrawal of some NSAIDs from the market, but symptomatic hepatic effects attributable to NSAIDs are rare and usually mild (Bannwarth and Berenbaum 2005). Lumiracoxib has a molecular phenyl acetic acid structure that is similar to that of diclofenac, the most widely prescribed NSAID world-wide, probably with an elevated risk for hepatotoxicity (which is estimated to occur in 4% of all patients). In the TARGET study the incidence of serious liver abnormalities (not further specified in the manuscript) was similar between the treatment groups, but lumiracoxib was associated with a RR of 3.97 (CI: 2.96–5.32) for increase in liver tests above 3 times the upper normal limit (Farkouh et al 2004; Schnitzer et al 2004). Thus, the risk of hepatotoxicity with the use of lumiracoxib is higher than for naproxen and ibuprofen. At the time of first submission of this manuscript, lumiracoxib was withdrawn from the market in Australia (where it had been on the market for several years and prescribed to 60,000 patients in doses up to 400 mg/day) because of 8 reports of serious liver adverse reactions to the drug, including 2 deaths and 2 liver transplants, further details of which were not yet available. Meanwhile, due to these serious liver adverse affects at doses >100 mg/day, lumiracoxib was withdrawn from the market in several countries, including the UK and Germany. In this context lumiracoxib should, were available, be limited to the lowest effective dose for the shortest possible duration of treatment. The reader is advised to inspect the safety issues that

<table>
<thead>
<tr>
<th>Study (reference)</th>
<th>Drugs</th>
<th>Side effects</th>
<th>Blood pressure</th>
<th>Hypertension</th>
<th>CHF</th>
<th>Hepatic</th>
<th>Cutaneous</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Renal</td>
<td>Edema</td>
<td>systolic</td>
<td>diastolic</td>
<td>opathy</td>
<td></td>
</tr>
<tr>
<td>VIGOR (Bombardier 2000)</td>
<td>Rofecoxib</td>
<td>1.2%</td>
<td>nr</td>
<td>nr</td>
<td>nr</td>
<td>nr</td>
<td>nr</td>
</tr>
<tr>
<td></td>
<td>Naproxen</td>
<td>0.9%</td>
<td>nr</td>
<td>nr</td>
<td>nr</td>
<td>nr</td>
<td>nr</td>
</tr>
<tr>
<td>CLASS (Silverstein 2000)</td>
<td>Celecoxib</td>
<td>0.7%*</td>
<td>2.8%</td>
<td>nr</td>
<td>nr</td>
<td>1.7%*</td>
<td>nr</td>
</tr>
<tr>
<td></td>
<td>Diclofenac</td>
<td>1.2%</td>
<td>3.5%</td>
<td>nr</td>
<td>nr</td>
<td>2.3%</td>
<td>nr</td>
</tr>
<tr>
<td>MEDAL (Cannon 2006; Laine 2007)</td>
<td>Etoricoxib</td>
<td>0.4%–2.3%</td>
<td>0.8%–1.9% (at 90 mg)</td>
<td>nr</td>
<td>nr</td>
<td>2.2%–2.5% **</td>
<td>0.1%–0.7%</td>
</tr>
<tr>
<td>TARGET (Schnitzer 2004; Farkouh 2004)</td>
<td>Lumiracoxib</td>
<td>***0.51%</td>
<td>0.4%–0.8%</td>
<td>+0.4 mm Hg to +0.1 mm Hg</td>
<td>+0.1 mm Hg</td>
<td>0.7%–1.6%</td>
<td>0.1%–0.3%</td>
</tr>
<tr>
<td></td>
<td>Naproxen/ Diclofenac</td>
<td>0.37%</td>
<td>+2.1 mm</td>
<td>+0.5 mm</td>
<td>+0.7%</td>
<td>0.07%</td>
<td>2.57%</td>
</tr>
</tbody>
</table>

*p < 0.05, **p < 0.001 vs control drug.
***p < 0.05 for discontinuations between treatment groups.
****major renal events, NS.
******serious liver abnormalities (not further specified), NS.
*******transaminases >> 3 times upper normal limit, p < 0.0001.

**Abbreviations:** ALT+, elevated alanine aminotransferase; AST+, elevated aspartate aminotransferase; CHF, cardiac heart failure; nr, not reported.
are emerging about liver safety, including baseline evaluation of history of liver diseases and regular checking liver function during treatment, and instructions about follow-up of liver function tests during treatment.

Conclusions

Lumiracoxib is a COX2 inhibitor that is highly selective, is more effective than placebo on pain in OA, with similar analgesic and anti-inflammatory effects as non-selective NSAIDs and the selective COX2 inhibitor celecoxib, has a lower incidence of upper GI side effects in patients not taking aspirin and a similar incidence of CV side effects as compared to naproxen or ibuprofen.

In the context of earlier guidelines (ACR Recommendations 2000; Hochberg 2002; Jordan et al 2003; Schnitzer 2006) and taking into account the GI and CV safety results of the TARGET study (Farkouh et al 2004; Schnitzer et al 2004), lumiracoxib is strictu sensu indicated in the treatment of patients with clinical OA of hip, knee or hands, or with radiographic OA of the spine, who do not respond to conventional treatment (such as analgesics [acetaminophen], physical therapy, and weight reduction in case of hip and knee OA), who have a moderate or high GI risk (with the restriction that in the TARGET study patients with a recent ulcer or bleeding or any history of perforation or obstruction were excluded) and a low CV risk and are not taking low dose aspirin (Table 5) (Chan 2006).

Lumiracoxib has been secured EMEA approval under the name of Prexige™ and Prexigen™ and has been launched in the UK since January 2006, where it is indicated for symptomatic treatment of osteoarthritis as well as short-term management of acute pain associated with primary dysmenorrhea and following orthopedic or dental surgery (www.emea.eu). The UK acted as the reference state in the EU’s mutual recognition procedure.

In the complex clinical context of efficiency and safety of selective and non-selective COX inhibitors, the prescription and use of COX2 inhibitors should be based on the risk and safety profile of the patient. One example is given in Table 5, in which the use of selective COX2 is proposed to be limited to patients with a low CV risk together with a moderate or high GI risk (Chan 2006). In addition, there is further need for long-term GI and CV safety studies on the use of selective and non-selective COX inhibitors. In view of the liver adverse effects, lumiracoxib should be limited to the lowest effective dose for the shortest possible duration of treatment, with special attention for liver toxicity according to the upcoming safety instructions. However, at the time of submission, lumiracoxib had been withdrawn from the market in several countries, including the UK and Germany, because of liver side effects at doses >100 mg/day. Meanwhile, at the time of proof approval of this paper, the European Medicines Agency (EMEA) had completed a review of the safety of medicines containing lumiracoxib. The Agency’s Committee for Medicinal Products for Human Use (CHMP) concluded that the benefits of these medicines no longer outweigh their risks, and that all marketing authorizations should be withdrawn throughout Europe because of the risk of serious side effects affecting the liver (www.emea.europa.eu).

References


