WTO’s new rules allow poorest to import drugs

The World Trade Organization is planning to put into law a waiver originally drafted in 2003 that allows least-developed countries to import generic drugs in public health emergencies, such as the HIV/AIDS epidemic.

The waiver pertains to Article 31 of the WTO’s Agreement on Trade-Related Aspects of Intellectual Property Rights (TRIPS), which requires that drugs manufactured under compulsory licensing be sold predominantly in the domestic market of the countries that produce them. If it stood, that provision would restrict the ability of countries lacking production capacity to import cheaper generic copies of patented drugs.

The amendment to Article 31 will come into force after ratification by two-thirds of WTO member countries — likely by December 2007. A statement attached to the agreement says that members will use the provision in “good faith” to deal with public health problems, not to meet industrial or commercial policy objectives.

The amendment “confirms once again that members are determined to ensure that WTO’s trading system contributes to humanitarian and development goals,” WTO Director-General Pascal Lamy stated in a news release.

But Médecins Sans Frontières says the 2003 waiver is cumbersome and inefficient: it requires each country to notify the WTO and to license the manufacture of generic drugs on a case-by-case basis. Once a generic manufacturer obtains a license, it can produce only enough medication to supply a single country. There is no provision in the agreement that would allow international tendering to procure the medicines, “which is the most common and efficient way of purchasing drugs,” says Carol Devine, acting program director for MSF.

So far, not one patient has benefited from the waiver, says Devine, who considers it “premature” to give it the force of law.

The International Federation of Pharmaceutical Manufacturers & Associations supports the amendment, saying it meets the needs of least-developed countries while preserving the agreement itself. — Laura Eggertson, CMAJ

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The vexations of Vioxx

The wave of litigation surrounding rofecoxib (Vioxx) has lapped ashore at the New England Journal of Medicine (NEJM), whose oft-cited study of the drug published in 2000 appears to contain incomplete adverse event information.

Merck & Co. pulled rofecoxib off the market on Sept. 30, 2004, after finding it doubled the risk of heart attack and stroke. There are now some 9200 lawsuits pending in the US against Merck & Co, based in Whitehouse Station, NJ.

In the midst of one such suit, on Nov. 21, 2005, a Merck & Co. memo dated July 5, 2000, emerged. It indicated that at least 2 authors of the NEJM article on the VIGOR (Vioxx Gastrointestinal Outcomes Research) study (2000;343:1520-8) knew 4.5 months before publication that 20 study participants suffered myocardial infarction (MI) after taking the drug — not 17 patients, as the NEJM article reported.

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more information about the safety and efficacy of the prostheses and supply a larger potential patient pool for conducting research, the report notes.

Inamed Corporation could not be reached for comment by press time, but a spokesman for Mentor Corporation issued the following statement about the panel report: “We believe in the scientific evidence supporting our products and look forward to Mentor’s Memory Gel breast implants becoming an important additional option for women seeking breast reconstruction or augmentation in Canada.”

Health Canada regulators will consider the panel report along with public comments and data from the manufacturer supporting its bid to get the products back on the Canadian market. A decision is expected in “a matter of months.”

Dr. Supriya Sharma, associate director general of Health Canada’s Therapeutic Products Directorate, said the recommendation for additional studies with regard to silicone bleed may affect the timing of any regulatory decision.

The report also recommended:

• Additional education for surgeons, since proper use is “crucial.” The panel “very strongly recommends” that Health Canada provide these prostheses only to certified Royal College plastic surgeons who have been specifically trained in implanting them.

• Patient information should acknowledge that the implant is not a lifetime device and will likely need to be replaced, necessitating subsequent surgery.

• Labelling that is available before surgery as printed material and on a Web site. Contraindications should include clinical depression, eating disorders and desire to breast feed.

• Patient and physician information should advise that multiple surgical procedures on the breast may cause irreversible changes to the breast itself, and physicians should be advised that “strong consideration should be given to implant removal” in the case of multiple surgeries. — Laura Eggertson, CMAJ

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