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This English version is intended to be a reference material to provide convenience for users. In the event of inconsistency between the Japanese original and this

English translation, the former shall prevail.

Revision of Precautions

Ibuprofen, celecoxib, naproxen, pranoprofen (oral dosage form), flurbiprofen axetil, loxoprofen sodium hydrate (oral dosage form), lornoxicam

February 25, 2021

Therapeutic category Antipyretics, analgesics and anti-inflammatory agents

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Non-proprietary name Ibuprofen, celecoxib, naproxen, pranoprofen, flurbiprofen axetil, loxoprofen sodium hydrate, lornoxicam

Safety measure

Precautions should be revised in the package insert.

Pharmaceuticals and Medical Devices Agency

3-3-2 Kasumigaseki, Chiyoda-ku, Tokyo 100-0013 Japan E-mail: <u>safety.info@pmda.go.jp</u> Revision in line with the Instructions for Package Inserts of Prescription Drugs, PAB Notification No. 606 by the Director General ofPharmaceutical Affairs Bureau, MHW, dated April 25, 1997 (Old instructions):Revised language is underlined.

Current	Revision
Use during Pregnancy, Delivery or Lactation	Use during Pregnancy, Delivery or Lactation
This drug should be administered to pregnant women (excluding	This drug should be administered to pregnant women (excluding
those in their third trimester) or women who may be pregnant only	those in their third trimester) or women who may be pregnant only
when the therapeutic benefits are considered to outweigh the risks.	when the therapeutic benefits are considered to outweigh the risks.
The safety of this drug administered during pregnancy has not	If such administration is deemed necessary, caution should be
been established.	exercised such as limiting to the minimum effective use and
	monitoring amniotic fluid as necessary. The safety of this drug
	administered during pregnancy has not been established. <u>Renal</u>
	impairment and decreased urine output in foetuses as well as
	accompanying oligohydramnios have been reported following use
	of cyclooxygenase inhibitors (oral dosage form or suppository) in
	pregnant women.

Revision in line with the Instructions for Package Inserts of Prescription Drugs, etc. PSEHB Notification No. 0608-1 by the Director of

Pharmaceutical Safety and Environmental Health Bureau, MHLW, dated June 8, 2017 (New instructions): Revised language is underlined.

Current	Revision
9. PRECAUTIONS CONCERNING PATIENTS WITH SPECIFIC	9. PRECAUTIONS CONCERNING PATIENTS WITH SPECIFIC
BACKGROUNDS	BACKGROUNDS
9.5 Pregnant Women	9.5 Pregnant Women
Pregnant women (excluding those in their third trimester) or women	Pregnant women (excluding those in their third trimester) or women
who may be pregnant	who may be pregnant

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This drug should be administered only when the therapeutic	This drug should be administered only when the therapeutic
benefits are considered to outweigh the risks.	benefits are considered to outweigh the risks. If such administration
	is deemed necessary, caution should be exercised such as limiting
	to the minimum effective use and monitoring amniotic fluid as
	necessary. Renal impairment and decreased urine output in
	foetuses as well as accompanying oligohydramnios have been
	reported following use of cyclooxygenase inhibitors (oral dosage
	form or suppository) in pregnant women.

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