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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 Talipexole hydrochloride

August 22, 2019

Therapeutic category

Antiparkinsonian agents

Non-proprietary name

Talipexole hydrochloride

Safety measure

Precautions should be revised in the package insert.

Pharmaceuticals and Medical Devices Agency

Revision in line with the Instructions for Package Inserts of Prescription Drugs, PAB Notification No. 606 by the Director General of Pharmaceutical Affairs Bureau, MHW, dated April 25, 1997 (Old instructions):

Revised language is underlined.

Current	Revision
Important Precautions	Important Precautions
When dose reduction or discontinuation of this drug is necessary,	When dose reduction or discontinuation of this drug is necessary,
the dose should be gradually reduced.	the dose should be gradually reduced. Rapid dose reduction or
	discontinuation may cause syndrome malin with symptoms such as
	pyrexia, disturbed consciousness, akinetic mutism, severe muscle
	stiffness, involuntary movement, dysphagia, tachycardia, blood
	pressure fluctuation, sweating, increased serum CK (CPK). In
	addition, rapid dose reduction or discontinuation of dopamine
	receptor agonists may cause drug withdrawal syndrome
	(characterized by apathy, anxiety, depression, fatigue, sweating,
	pain, etc.).
Adverse Reactions	Adverse Reactions
Other Adverse Reactions	Other Adverse Reactions
(N/A)	Drug withdrawal syndrome* (apathy, anxiety, depression, fatigue,
	sweating, pain, etc.)
	*When any abnormalities are observed, appropriate measures
	should be taken such as resuming administration or returning the
	dose to the level prior to reduction.

N/A: Not Applicable, because the section is not included in the current package insert.