

**IMPLEMENTATION OF THE VARIATION REGULATION N° 1234/2008 IN EACH MEMBER
STATE FOR MEDICINAL PRODUCTS AUTHORISED BY PURELY NATIONAL PROCEDURES**

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It is reminded that It is not possible to combine nationally and European authorised products in the same variation until the Variation Regulation is amended allowing to apply the provisions to purely national products.

Will the variation regulation N° 1234/2008 be implemented from 1st January 2010 also for medicinal products authorised by purely national procedures?

	AT	BE	BG	CY	CZ	DE	DK	EE	EL	ES	FI	FR	HU	IE	IS	IT	LI	LT	LU	LV	MT	NL	NO	PL	PT	RO	SE	SI	SK	UK	
No	X		X		X	X						X								X				X	X						
Yes		X		X				X	X				X			X						X						X	X	X	
Yes, but different timelines							X			X	X			X	X			X		X						X	X				