

Message 001

Communication from the Commission - TRIS/(2025) 1508

Directive (EU) 2015/1535

Notification: 2025/0294/FR

Notification of a draft text from a Member State

Notification – Notification – Notifizierung – Нотификация – Oznámení – Notifikation – Γνωστοποίηση – Notificación – Teavitamine – Ilmoitus – Obavijest – Bejelentés – Notifica – Pranešimas – Paziņojums – Notifika – Kennisgeving – Zawiadomienie – Notificação – Notificare – Oznámenie – Obvestilo – Anmälan – Fógra a thabhairt

Does not open the delays - N'ouvre pas de délai - Kein Fristbeginn - Не се предвижда период на прекъсване - Nezahtuje prodlení - Fristerne indledes ikke - Καμμία έναρξη προθεσμίας - No abre el plazo - Viivituste perioodi ei avata - Määräaika ei ala tästä - Ne otvara razdoblje kašnjenja - Nem nyitja meg a késések - Non fa decorrere la mora - Atidėjimai nepradedami - Atlikšanas laikposms nesākas - Ma jiftaħ il-perijodi ta' dewmien - Geen termijnbegin - Nie otwiera opóźnień - Não inicia o prazo - Nu deschide perioadele de stagnare - Nezačína oneskorenia - Ne uvaja zamud - Inleder ingen frist - Ní osclaíonn sé na moilleanna

MSG: 20251508.EN

1. MSG 001 IND 2025 0294 FR EN 10-06-2025 FR NOTIF

2. France

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3B. Agence nationale de sécurité du médicament et des produits de santé –

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4. 2025/0294/FR - C30P - Pharmacopoeias

5. Pro Pharmacopoeia technical note submitted for public consultation No 1294 – Ambra grisea

6. Draft monographs for pharmaceutical products

7.

8. It is proposed to revise this monograph to improve techniques or specifications related to the quality of the substances authorised on the market.

9. The texts, implemented in the 11th edition of the French Pharmacopoeia, will be published by decision of the Director-General on the National Agency for the Safety of Medicines and Health Products website, with the following statement: 'Having regard to Directive (EU) 2015/1535 of the European Parliament and of the Council of 9 September 2015 laying down a procedure for the provision of information in the field of technical regulations and of rules on Information Society services, and in particular Notification No ...'

10. References to normative texts: There are no normative texts

11. No

12.

13. No

14. No

15. No

16.

TBT aspects: No

SPS aspects: No

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European Commission

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