



EUROPEAN MEDICINES AGENCY  
SCIENCE MEDICINES HEALTH

10 July 2025  
EMADOC-1700519818-2288925  
Human Medicines Division

## List of nationally authorised medicinal products

Active substance(s): acetylsalicylic acid / caffeine / paracetamol

EURD list No. PSUSA/00002291/202412



| <b>Product Name (in authorisation country)</b>                        | <b>MRP/DCP Authorisation number</b> | <b>National Authorisation Number</b> | <b>MAH of product in the member state</b> | <b>Member State where product is authorised</b> |
|-----------------------------------------------------------------------|-------------------------------------|--------------------------------------|-------------------------------------------|-------------------------------------------------|
| Actron Compuesto 267 mg / 133 mg / 40 mg comprimidos efervescentes    | not available                       | 47.178                               | BAYER HISPANIA SL                         | ES                                              |
| ACTRON, comprimé effervescent                                         | not available                       | 34009 311 834 0 2                    | BAYER HEALTHCARE                          | FR                                              |
| ACTRON, comprimé effervescent                                         | not available                       | 34009 340 211 8 3                    | BAYER HEALTHCARE                          | FR                                              |
| ACTRON, comprimé effervescent                                         | not available                       | 34009 336 991 2 3                    | BAYER HEALTHCARE                          | FR                                              |
| ACTRON, comprimé effervescent                                         | not available                       | 34009 322 138 0 1                    | BAYER HEALTHCARE                          | FR                                              |
| Algon<br>Δισκία (400+200+50) mg/tab                                   | not available                       | 3151301                              | LAVIPHARM S.A                             | GR                                              |
| Alka-Seltzer XS                                                       | not available                       | PL 00010/0510                        | BAYER PLC                                 | XI                                              |
| Anadin Extra                                                          | not available                       | PL 44673/0199                        | HALEON UK TRADING LIMITED                 | XI                                              |
| ANTIREUMINA                                                           | not available                       | 004172021                            | TEOFARMA S.R.L.                           | IT                                              |
| Excedite 250 mg / 250 mg / 65 mg comprimidos recubiertos con película | DE/H/1494/001                       | 72543                                | HALEON SPAIN, S.A.                        | ES                                              |
| Excedrin 250 mg /250 mg /65 mg film-coated tablets                    | DE/H/1494/001                       | PA0678/122/001                       | HALEON IRELAND LIMITED                    | IE                                              |
| Excedrin MigraStop, 250 mg + 250 mg + 65 mg, tabletki powlekane       | DE/H/1494/001                       | 16490                                | HALEON POLAND SP. Z O.O.                  | PL                                              |
| EXCEDRIN, filmomhulde tabletten                                       | DE/H/1494/001                       | RVG 102780                           | HALEON NETHERLANDS B.V.                   | NL                                              |
| Excedryn Migraine & Hoofdpijn, filmomhulde tabletten                  | DE/H/1494/001                       | BE352493                             | HALEON BELGIUM                            | BE                                              |

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| Excedryn Migraine & Maux de tête, comprimés pelliculés                | DE/H/1494/001                       | BE352493                             | HALEON BELGIUM                              | BE                                              |
| Excedryn Migraine & Maux de tête, comprimés pelliculés                | DE/H/1494/001                       | 2010040086                           | HALEON BELGIUM                              | LU                                              |
| EXCEDRYN Migräne & Kopfschmerzen, Filmtabletten                       | DE/H/1494/001                       | BE352493                             | HALEON BELGIUM                              | BE                                              |
| EXCEDRYN Migräne & Kopfschmerzen, Filmtabletten                       | DE/H/1494/001                       | 2010040086                           | HALEON BELGIUM                              | LU                                              |
| Panadol Migrena 250 mg/250 mg/65 mg filmom obalene tablety            | DE/H/1494/001                       | 07/0555/09-S                         | HALEON CZECH REPUBLIC S.R.O.                | SK                                              |
| Panmigran 250 mg / 250 mg / 65 mg επικαλυμμένα με λεπτό υμένιο δισκία | DE/H/1494/001                       | 023953                               | HALEON HELLAS SINGLE MEMBER SOCIETE ANONYME | CY                                              |
| Panmigran 250 mg / 250 mg / 65 mg επικαλυμμένα με λεπτό υμένιο δισκία | DE/H/1494/001                       | 319400111                            | HALEON HELLAS SINGLE MEMBER SOCIETE ANONYME | GR                                              |
| Panmigran 250 mg / 250 mg / 65 mg επικαλυμμένα με λεπτό υμένιο δισκία | DE/H/1494/001                       | 319400113                            | HALEON HELLAS SINGLE MEMBER SOCIETE ANONYME | GR                                              |
| Panmigran 250 mg / 250 mg / 65 mg επικαλυμμένα με λεπτό υμένιο δισκία | DE/H/1494/001                       | 319400116                            | HALEON HELLAS SINGLE MEMBER SOCIETE ANONYME | GR                                              |
| Panmigran 250 mg / 250 mg / 65 mg επικαλυμμένα με λεπτό υμένιο δισκία | DE/H/1494/001                       | 319400110                            | HALEON HELLAS SINGLE MEMBER SOCIETE ANONYME | GR                                              |
| Panmigran 250 mg / 250 mg / 65 mg επικαλυμμένα με λεπτό υμένιο δισκία | DE/H/1494/001                       | 319400109                            | HALEON HELLAS SINGLE                        | GR                                              |

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| mg / 65 mg<br>επικαλυμμένα με λεπτό υμένιο δισκία                        |                                     |                                      | MEMBER SOCIETE ANONYME                      |                                                 |
| Panmigran 250 mg / 250 mg / 65 mg<br>επικαλυμμένα με λεπτό υμένιο δισκία | DE/H/1494/001                       | 319400114                            | HALEON HELLAS SINGLE MEMBER SOCIETE ANONYME | GR                                              |
| Panmigran 250 mg / 250 mg / 65 mg<br>επικαλυμμένα με λεπτό υμένιο δισκία | DE/H/1494/001                       | 319400115                            | HALEON HELLAS SINGLE MEMBER SOCIETE ANONYME | GR                                              |
| Panmigran 250 mg / 250 mg / 65 mg<br>επικαλυμμένα με λεπτό υμένιο δισκία | DE/H/1494/001                       | 319400112                            | HALEON HELLAS SINGLE MEMBER SOCIETE ANONYME | GR                                              |
| Panmigran 250 mg / 250 mg / 65 mg<br>επικαλυμμένα με λεπτό υμένιο δισκία | DE/H/1494/001                       | 319400101                            | HALEON HELLAS SINGLE MEMBER SOCIETE ANONYME | GR                                              |
| Panmigran 250 mg / 250 mg / 65 mg<br>επικαλυμμένα με λεπτό υμένιο δισκία | DE/H/1494/001                       | 319400102                            | HALEON HELLAS SINGLE MEMBER SOCIETE ANONYME | GR                                              |
| Panmigran 250 mg / 250 mg / 65 mg<br>επικαλυμμένα με λεπτό υμένιο δισκία | DE/H/1494/001                       | 319400106                            | HALEON HELLAS SINGLE MEMBER SOCIETE ANONYME | GR                                              |
| Panmigran 250 mg / 250 mg / 65 mg<br>επικαλυμμένα με λεπτό υμένιο δισκία | DE/H/1494/001                       | 319400103                            | HALEON HELLAS SINGLE MEMBER SOCIETE ANONYME | GR                                              |
| Panmigran 250 mg / 250 mg / 65 mg<br>επικαλυμμένα με λεπτό υμένιο δισκία | DE/H/1494/001                       | 319400107                            | HALEON HELLAS SINGLE MEMBER SOCIETE ANONYME | GR                                              |
| Panmigran 250 mg / 250                                                   | DE/H/1494/001                       | 319400108                            | HALEON HELLAS SINGLE                        | GR                                              |

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| mg / 65 mg<br>επικαλυμμένα με λεπτό υμένιο δισκία                                   |                                     |                                      | MEMBER SOCIETE ANONYME                                                   |                                                 |
| Panmigran 250 mg / 250 mg / 65 mg<br>επικαλυμμένα με λεπτό υμένιο δισκία            | DE/H/1494/001                       | 319400105                            | HALEON HELLAS SINGLE MEMBER SOCIETE ANONYME                              | GR                                              |
| Panmigran 250 mg / 250 mg / 65 mg<br>επικαλυμμένα με λεπτό υμένιο δισκία            | DE/H/1494/001                       | 319400104                            | HALEON HELLAS SINGLE MEMBER SOCIETE ANONYME                              | GR                                              |
| Panmigran 250 mg /250 mg /65 mg film-coated tablets                                 | DE/H/1494/001                       | MA1177/01301                         | GLAXOSMITHKLINE CONSUMER HEALTHCARE HELLAS SINGLE MEMBER SOCIETE ANONYME | MT                                              |
| Panmigrol Migraine and Headache Relief 250 mg /250 mg /65 mg film coated tablets    | not available                       | PL 44673/0229                        | HALEON UK TRADING LIMITED                                                | XI                                              |
| Panmigrol Migraine Pain Relief 250 mg /250 mg /65 mg film coated tablets            | not available                       | PL 44673/0231                        | HALEON UK TRADING LIMITED                                                | XI                                              |
| Paracetamol/ASS/Coffein Kopfschmerz & Migräne 250 mg / 250 mg / 65 mg Filmdabletten | DE/H/1494/001                       | 71852.00.00                          | HALEON GERMANY GMBH                                                      | DE                                              |
| Perdolan Compositum Suppositories                                                   | not available                       | 2002106930                           | JOHNSON & JOHNSON CONSUMER N.V./S.A.                                     | LU                                              |
| Perdolan Compositum Suppositories                                                   | not available                       | 2002106930                           | JOHNSON & JOHNSON CONSUMER N.V./S.A.                                     | LU                                              |
| Perdolan Compositum Suppositories                                                   | not available                       | 2002106930                           | JOHNSON & JOHNSON CONSUMER N.V./S.A.                                     | LU                                              |
| Perdolan Compositum Suppositories                                                   | not available                       | 2002106930                           | JOHNSON & JOHNSON CONSUMER N.V./S.A.                                     | LU                                              |
| PERDOLAN                                                                            | not available                       | BE228277                             | JOHNSON & JOHNSON                                                        | BE                                              |

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| COMPOSITUM suppositoires                                                 |                              |                               | CONSUMER N.V./S.A.                                       |                                          |
| PERDOLAN COMPOSITUM suppositoires                                        | not available                | BE228277                      | JOHNSON & JOHNSON CONSUMER N.V./S.A.                     | BE                                       |
| PERDOLAN COMPOSITUM Zäpfchen                                             | not available                | BE228277                      | JOHNSON & JOHNSON CONSUMER N.V./S.A.                     | BE                                       |
| PERDOLAN COMPOSITUM Zäpfchen                                             | not available                | BE228277                      | JOHNSON & JOHNSON CONSUMER N.V./S.A.                     | BE                                       |
| PERDOLAN COMPOSITUM zetpillen                                            | not available                | BE228277                      | JOHNSON & JOHNSON CONSUMER N.V./S.A.                     | BE                                       |
| PERDOLAN COMPOSITUM zetpillen                                            | not available                | BE228277                      | JOHNSON & JOHNSON CONSUMER N.V./S.A.                     | BE                                       |
| SPLENTIR (400+200+50) mg/tab<br>Αναβράζοντα δισκία                       | not available                | 1884003 (PRODUCT CODE)        | UNI-PHARMA KLEON TSETIS PHARMACEUTICAL LABORATORIES S.A. | GR                                       |
| SPLENTIR (400+200+50) mg/tab<br>Δισκία                                   | not available                | 1884001                       | UNI-PHARMA KLEON TSETIS PHARMACEUTICAL LABORATORIES S.A. | GR                                       |
| SPLENTIR (400+200+50) mg/tab<br>Επικαλυμμένα δισκία                      | not available                | 1884002                       | UNI-PHARMA KLEON TSETIS PHARMACEUTICAL LABORATORIES S.A. | GR                                       |
| Thomapyrin - Tabletten                                                   | not available                | 1-17100                       | OPELLA HEALTHCARE AUSTRIA GMBH                           | AT                                       |
| Thomapyrin - Tabletten                                                   | not available                | 1-17100                       | OPELLA HEALTHCARE AUSTRIA GMBH                           | AT                                       |
| Thomapyrin® CLASSIC Schmerztabletten<br>250 mg/200 mg/50 mg pro Tablette | not available                | 6134.00.00                    | A. NATTERMANN & CIE. GMBH                                | DE                                       |
| Thomapyrin® CLASSIC Schmerztabletten<br>250 mg/200 mg/50 mg pro Tablette | not available                | 6134.00.00                    | A. NATTERMANN & CIE. GMBH                                | DE                                       |
| Thomapyrin® CLASSIC Schmerztabletten                                     | not available                | 6134.00.00                    | A. NATTERMANN & CIE. GMBH                                | DE                                       |

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| 250 mg/200 mg/50 mg pro Tablette                        |                                     |                                      |                                           |                                                 |
| Thomapyrin® INTENSIV 250 mg/250 mg/50 mg pro Tablette   | not available                       | 6386585.00.00                        | A. NATTERMANN & CIE. GMBH                 | DE                                              |
| Voltamig 250 mg / 250 mg / 65 mg comprimidos revestidos | DE/H/1494/001                       | 5262209                              | HALEON PORTUGAL, LDA                      | PT                                              |
| Voltamig 250 mg / 250 mg / 65 mg comprimidos revestidos | DE/H/1494/001                       | 5262175                              | HALEON PORTUGAL, LDA                      | PT                                              |
| Voltamig 250 mg / 250 mg / 65 mg comprimidos revestidos | DE/H/1494/001                       | 5262217                              | HALEON PORTUGAL, LDA                      | PT                                              |