VPA10774/054/004

Selehold 240 mg spot-on solution for dogs 20.1–40.0 kg

| Variation | Summary | Date |
|------------------|---|----------|
| Vet - A1 e) | VNRA - Vet - A1 e) - e) Change in the name or address or contact details of a manufacturer or importer of the finished product (including batch release or quality control testing sites) - A1 e) Administrative changes: Change in the name or address or contact details of a manufacturer or importer of the finished product (including batch release or quality control testing sites) | 19/05/25 |
| Vet - F.I.a.1 z) | VRA-R - Vet - F.I.a.1 z) - z) Other changes under this code level e.g. variations outlined in section 6 and 7 of EMA/CMDv/7381/2021 - F.I.a.1 z) - Quality Changes - Active Substance - Manufacture - Change in the manufacturer of a starting material/reagent/intermediate used in the manufacturing process of the active substance or change in the manufacturer (including where relevant quality control testing sites) of the active substance, where no Ph. Eur. Certificate of Suitability is part of the approved dossier - Other changes under this code level, e.g. variations outlined in section 6 and 7 of EMA/CMDv/7381/2021 | 04/03/25 |
| Vet - B3 a) | VNRA - Vet - B3 a) - a) Deletion of a manufacturing site for an active substance, intermediate or finished product, packaging site, manufacturer responsible for batch release, site where batch control takes place, or supplier of a starting material for an active substance, reagent or excipient (when mentioned in the dossier) - B3 a) Changes to the quality part of the dossier: Deletion of a manufacturing site for an active substance, intermediate or finished product, packaging site, manufacturer responsible for batch release, site where batch control takes place, or supplier of a starting material for a starting material for a starting material for an active substance, intermediate or finished product, packaging site, manufacturer responsible for batch release, site where batch control takes place, or supplier of a starting material for an active substance, reagent or excipient (when mentioned in the dossier) | 21/11/24 |
| Vet - C6 | VNRA - Vet - C6 - Introduction of a summary of the PSMF or changes to the summary of the PSMF not already covered elsewhere in the Annex to Regulation (EU) 2021/17 - C6 Changes to the safety, efficacy and pharmacovigilance part of the dossier: Introduction of a summary of the PSMF or changes to the summary of the PSMF not already covered elsewhere in the Annex to Regulation (EU) 2021/17 | 18/10/24 |
| Vet - G.I.18 | VRA-S - Vet - G.I.18 - One-off alignment of the product information with version 9.0 (or the latest version of the QRD templates that are in effect at the time that this one-off variation is submitted) of the QRD templates i.e. major update of the QRD templates in accordance with Regulation (EU) 2019/6, for veterinary medicinal products placed on the market in accordance with Directive 2001/82/EC or Regulation (EC) No 726/2004 - G.I.18 Safety, Efficacy, Pharmacovigilance changes - One-off alignment of the product information with version 9.0 (or the | 15/09/23 |

| | latest version of the QRD templates that are in effect at the time | |
|------------------------------|--|----------|
| | that this one-off variation is submitted) of the QRD templates i.e. | |
| | major update of the QRD templates in accordance with Regulation (EU) 2019/6, for veterinary medicinal products placed | |
| | on the market in accordance with Directive 2001/82/EC or | |
| | | |
| | Regulation (EC) No 726/2004 VNRA - Vet - B12 c) - c) Minor changes to an approved test | |
| Vet - B12 c) Vet - B47 d) | procedure for an in-process test - B12 c) Changes to the quality | |
| | part of the dossier: Minor changes — to an approved test | 24/03/23 |
| | procedure for an in-process test — for active substance; — for the | 24/03/23 |
| | finished product | |
| | VNRA - Vet - B47 d) - d) To reflect compliance with the Ph. Eur. | |
| | by removing reference to the internal test method and test method | |
| | number - B47d) Changes to the quality part of the dossier: | |
| | Change to comply with Ph. Eur. or with a national pharmacopoeia | 02/03/23 |
| | of a Member State: — to reflect compliance with the Ph. Eur. by | 02/03/23 |
| | removing reference to the internal test method and test method | |
| | number | |
| | VNRA - Vet - B4 a) - a) Change in the manufacturer of the active | |
| | substance (including relevant quality control testing sites) - B4 a) | |
| | Changes to the quality part of the dossier: Changes to the | |
| Vet - B4 a) | production process or the storage of active substance where no | 00/02/02 |
| | Ph. Eur. CEP is part of the approved dossier of an active | 02/03/23 |
| | substance (including starting material, reagent or intermediate) - | |
| | change in the manufacturer of the active substance (including | |
| | relevant quality control testing sites) | |
| Vet - F.I.d.1 c) | VRA-R - Vet - F.I.d.1 c) - c) Extension or introduction of a re-test | |
| | period/storage period supported by real time data - F.I.d.1 c) | |
| | Quality Changes - Active Substance - Stability - Change in the | |
| | | 08/08/22 |
| | Eur. Certificate of Suitability covering the retest period is part of | |
| | the approved dossier - Extension or introduction of a re-test | |
| | period/storage period supported by real time data | |
| B.II.b.5.z | IA - B.II.b.5.z - z Other variation - B.II.b.5.z - QUALITY | |
| | CHANGES - FINISHED PRODUCT - Manufacture - Change to | 28/02/22 |
| | in-process tests or limits applied during the manufacture of the | |
| | finished product - Other variation | |
| B.II.d.2.a | IA - B.II.d.2.a - a) Minor changes to an approved test procedure - | 28/02/22 |
| | B.II.d.2.a - QUALITY CHANGES - FINISHED PRODUCT - | |
| | Control of finished product - Change in test procedure for the | |
| | finished product - Minor changes to an approved test procedure | |