

## Ozurdex

Procedural steps taken and scientific information after the authorisation

| Application<br>number | Scope   | Opinion/<br>Notification<br><sup>1</sup> issued on | Commission<br>Decision<br>Issued <sup>2</sup> /<br>amended<br>on | Product<br>Information<br>affected <sup>3</sup> | Summary |
|-----------------------|---|--|--|---|---------|
| II/0044               | Submission of an updated RMP version 12.1.<br>C.I.11.b - Introduction of, or change(s) to, the<br>obligations and conditions of a marketing<br>authorisation, including the RMP - Implementation of<br>change(s) which require to be further substantiated<br>by new additional data to be submitted by the MAH | 26/10/2023   | n/a  |   |         |

<sup>&</sup>lt;sup>1</sup> Notifications are issued for type I variations and Article 61(3) notifications (unless part of a group including a type II variation or extension application or a worksharing application). Opinions are issued for all other procedures.

- <sup>2</sup> A Commission decision (CD) is issued for procedures that affect the terms of the marketing authorisation (e.g. summary of product characteristics, annex II, labelling, package leaflet). The
- CD is issued within two months of the opinion for variations falling under the scope of Article 23.1a(a) of Regulation (EU) No. 712/2012, or within one year for other procedures.



<sup>&</sup>lt;sup>3</sup> SmPC (Summary of Product Characteristics), Annex II, Labelling, PL (Package Leaflet).

|                      | where significant assessment is required   |            |            |                              |                                   |
|----------------------|--|------------|------------|------------------------------|-----------------------------------|
| PSUSA/985/2<br>02201 | Periodic Safety Update EU Single assessment -<br>dexamethasone (centrally authorised product<br>indicated in uveitis and macular oedema)   | 29/09/2022 | n/a        |                              | PRAC Recommendation - maintenance |
| N/0042               | Minor change in labelling or package leaflet not connected with the SPC (Art. 61.3 Notification)   | 20/07/2022 |            | PL                           |                                   |
| T/0040               | Transfer of Marketing Authorisation  | 29/04/2022 | 13/06/2022 | SmPC,<br>Labelling and<br>PL |                                   |
| N/0039               | Minor change in labelling or package leaflet not connected with the SPC (Art. 61.3 Notification)   | 23/06/2021 | 13/06/2022 | PL                           |                                   |
| N/0038               | Minor change in labelling or package leaflet not connected with the SPC (Art. 61.3 Notification)   | 26/02/2021 | 13/06/2022 | PL                           |                                   |
| 11/0037              | Submission of an updated RMP version 9.0 (and 10.0<br>during the procedure) in order to reflect increased<br>knowledge of the product and align to the new RMP<br>template.<br>C.I.11.b - Introduction of, or change(s) to, the<br>obligations and conditions of a marketing<br>authorisation, including the RMP - Implementation of<br>change(s) which require to be further substantiated<br>by new additional data to be submitted by the MAH<br>where significant assessment is required | 17/04/2020 | n/a        |                              |                                   |

| PSUSA/985/2<br>01901 | Periodic Safety Update EU Single assessment -<br>dexamethasone (centrally authorised product<br>indicated in uveitis and macular oedema)   | 05/09/2019 | n/a        |                          | PRAC Recommendation - maintenance   |
|----------------------|--|------------|------------|--------------------------|---|
| 11/0035              | Submission of the final report from study CMO-EPI-<br>EYE-0522 listed as a category 3 study in the RMP.<br>This is an observational, cross-sectional study<br>conducted in France, Germany, Spain, and<br>the UK having as primary objective the assessment<br>of the effectiveness of the educational material<br>provided to the treating physicians.<br>The SmPC sections 4.2, 6.6 and Annex II of Product<br>information were updated to reflect the conclusions<br>of the assessment. Package Leaflet is updated<br>accordingly.<br>In addition, the Marketing authorisation holder took<br>the opportunity to include updates to local<br>representative in SK.<br>C.I.13 - Other variations not specifically covered<br>elsewhere in this Annex which involve the submission<br>of studies to the competent authority | 05/09/2019 |            | SmPC, Annex<br>II and PL | To continue providing physicians the intravitreal injection<br>procedure pictogram, the illustrations are included in the<br>product information. In SmPC section 4.2 a cross-reference<br>to section 6.6 in which the intravitreal injection procedure<br>pictogram is included. Information in the package leaflet is<br>included under section 6.6 of the "Information for the<br>healthcare professional" provided in the carton. |
| IB/0034              | B.II.d.2.d - Change in test procedure for the finished<br>product - Other changes to a test procedure<br>(including replacement or addition)   | 28/05/2019 | n/a        |                          |   |
| II/0032              | C.I.3.b - Change(s) in the SPC, Labelling or PL<br>intended to implement the outcome of a procedure<br>concerning PSUR or PASS or the outcome of the<br>assessment done under A 45/46 - Change(s) with   | 28/03/2019 | 10/03/2020 | SmPC and PL              |   |

|           | new additional data submitted by the MAH               |            |            |    |
|-----------|--|------------|------------|----|
| N/0033    | Minor change in labelling or package leaflet not       | 22/03/2019 | 10/03/2020 | PL |
|           | connected with the SPC (Art. 61.3 Notification)        |            |            |    |
|           |  |            |            |    |
| IB/0031/G | This was an application for a group of variations.     | 05/02/2019 | n/a        |    |
|           |  |            |            |    |
|           | B.II.b.5.a - Change to in-process tests or limits      |            |            |    |
|           | applied during the manufacture of the finished         |            |            |    |
|           | product - Tightening of in-process limits              |            |            |    |
|           | B.II.b.5.b - Change to in-process tests or limits      |            |            |    |
|           | applied during the manufacture of the finished         |            |            |    |
|           | product - Addition of a new test(s) and limits         |            |            |    |
|           | B.II.b.5.f - Change to in-process tests or limits      |            |            |    |
|           | applied during the manufacture of the finished         |            |            |    |
|           | product - Addition or replacement of an in-process     |            |            |    |
|           | test as a result of a safety or quality issue          |            |            |    |
|           | B.II.b.5.f - Change to in-process tests or limits      |            |            |    |
|           | applied during the manufacture of the finished         |            |            |    |
|           | product - Addition or replacement of an in-process     |            |            |    |
|           | test as a result of a safety or quality issue          |            |            |    |
|           | B.II.b.5.z - Change to in-process tests or limits      |            |            |    |
|           | applied during the manufacture of the finished         |            |            |    |
|           | product - Other variation                              |            |            |    |
|           | B.II.d.1.a - Change in the specification parameters    |            |            |    |
|           | and/or limits of the finished product - Tightening of  |            |            |    |
|           | specification limits                                   |            |            |    |
|           | B.II.d.1.a - Change in the specification parameters    |            |            |    |
|           | and/or limits of the finished product - Tightening of  |            |            |    |
|           | specification limits                                   |            |            |    |
|           | B.II.d.2.d - Change in test procedure for the finished |            |            |    |

|                      | product - Other changes to a test procedure<br>(including replacement or addition)<br>B.II.d.2.d - Change in test procedure for the finished<br>product - Other changes to a test procedure<br>(including replacement or addition) |            |            |                     |  |
|----------------------|--|------------|------------|---------------------|--|
| PSUSA/985/2<br>01801 | Periodic Safety Update EU Single assessment -<br>dexamethasone (centrally authorised product<br>indicated in uveitis and macular oedema)   | 20/09/2018 | 20/11/2018 | SmPC                | Refer to Scientific conclusions and grounds recommending<br>the variation to terms of the Marketing Authorisation(s)' for<br>PSUSA/985/201801. |
| IB/0030              | B.II.b.4.a - Change in the batch size (including batch<br>size ranges) of the finished product - Up to 10-fold<br>compared to the originally approved batch size   | 28/08/2018 | n/a        |                     |  |
| N/0028               | Minor change in labelling or package leaflet not connected with the SPC (Art. 61.3 Notification)   | 06/04/2018 | 20/11/2018 | Labelling and<br>PL |  |
| PSUSA/985/2<br>01701 | Periodic Safety Update EU Single assessment -<br>dexamethasone (centrally authorised product<br>indicated in uveitis and macular oedema)   | 14/09/2017 | 15/11/2017 | SmPC                | Refer to Scientific conclusions and grounds recommending<br>the variation to terms of the Marketing Authorisation(s)' for<br>PSUSA/985/201701. |
| IA/0027              | B.III.1.a.2 - Submission of a new/updated or<br>deletion of Ph. Eur. Certificate of Suitability to the<br>relevant Ph. Eur. Monograph - Updated certificate<br>from an already approved manufacturer                               | 01/06/2017 | n/a        |                     |  |
| II/0025              | In line with the RMP commitment, submission of the<br>final report for the Post-Authorisation Safety Study<br>206207-025 (A Prospective Observational Study to<br>Evaluate Long-Term Safety in Real-World Clinical<br>Practice).   | 23/03/2017 | n/a        |                     |  |

|                      | C.I.13 - Other variations not specifically covered<br>elsewhere in this Annex which involve the submission<br>of studies to the competent authority   |            |            |    |                                   |
|----------------------|---|------------|------------|----|-----------------------------------|
| PSUSA/985/2<br>01601 | Periodic Safety Update EU Single assessment -<br>dexamethasone (centrally authorised product<br>indicated in uveitis and macular oedema)  | 02/09/2016 | n/a        |    | PRAC Recommendation - maintenance |
| IB/0022              | C.I.11.z - Introduction of, or change(s) to, the obligations and conditions of a marketing authorisation, including the RMP - Other variation   | 24/02/2016 | n/a        |    |                                   |
| N/0023               | Update of the package leaflet with revised contact<br>details of the local representatives. The MAH also<br>took the opportunity to update section 4 of the<br>package leaflet to add missing asterisks for ADRs<br>that are related to the injection procedure, in line<br>with section 4.8 of the SmPC ADRs.<br>Minor change in labelling or package leaflet not<br>connected with the SPC (Art. 61.3 Notification) | 23/02/2016 | 15/11/2017 | PL |                                   |
| IA/0021              | B.III.1.a.2 - Submission of a new/updated or<br>deletion of Ph. Eur. Certificate of Suitability to the<br>relevant Ph. Eur. Monograph - Updated certificate<br>from an already approved manufacturer  | 28/01/2016 | n/a        |    |                                   |
| PSUSA/985/2<br>01501 | Periodic Safety Update EU Single assessment -<br>dexamethasone (centrally authorised product<br>indicated in uveitis and macular oedema)  | 10/09/2015 | n/a        |    | PRAC Recommendation - maintenance |

| IAIN/0020 | C.I.8.a - Introduction of or changes to a summary of<br>Pharmacovigilance system - Changes in QPPV<br>(including contact details) and/or changes in the<br>PSMF location  | 21/05/2015 | n/a        |  |   |
|-----------|---|------------|------------|--|---|
| R/0018    | Renewal of the marketing authorisation.   | 22/01/2015 | 23/03/2015 | SmPC, Annex<br>II, Labelling<br>and PL | Based on the review of available information, the CHMP is<br>of the opinion that the quality, safety and efficacy of<br>Ozurdex continues to be adequately and sufficiently<br>demonstrated and considers that the benefit/risk profile of<br>this medicinal product continues to be favourable.<br>The product information has been updated to align with<br>QRD templates.<br>The CHMP recommends that the renewal be granted with<br>unlimited validity. |
| PSUV/0017 | Periodic Safety Update  | 11/09/2014 | n/a        |  | PRAC Recommendation - maintenance   |
| II/0015   | Extension of Indication to include a new indication<br>for the treatment of adult patients with visual<br>impairment due to diabetic macular oedema who are<br>pseudophakic or who are considered insufficiently<br>responsive to, or unsuitable for non-corticosteroid<br>therapy.<br>C.I.6.a - Change(s) to therapeutic indication(s) -<br>Addition of a new therapeutic indication or<br>modification of an approved one | 24/07/2014 | 26/08/2014 | SmPC, Annex<br>II and PL               | Please refer to scientific summary.   |
| IA/0016/G | This was an application for a group of variations.  | 21/05/2014 | n/a        |  |   |
|           | B.II.e.7.a - Change in supplier of packaging  |            |            |  |   |

|         | components or devices (when mentioned in the<br>dossier) - Deletion of a supplier<br>B.II.a.3.b.1 - Changes in the composition<br>(excipients) of the finished product - Other excipients<br>- Any minor adjustment of the quantitative<br>composition of the finished product with respect to<br>excipients<br>B.II.e.3.c - Change in test procedure for the<br>immediate packaging of the finished product -<br>Deletion of a test procedure if an alternative test<br>procedure is already authorised  |            |            |                    |   |
|---------|---|------------|------------|--------------------|---|
| II/0013 | Update of Annex II to remove reference to the<br>injection procedure video from the list of elements of<br>the educational material package in the conditions<br>and restrictions for the safe and effective use of the<br>medicinal product. Additional updates to the Risk<br>Management Plan (RMP) were made to consolidate<br>the information in the educational material as well as<br>in response to previous requests following the<br>assessment of RMP version 1.9 and PSUR #4.<br>C.I.z - Changes (Safety/Efficacy) of Human and<br>Veterinary Medicinal Products - Other variation | 27/06/2013 | 19/06/2014 | Annex II and<br>PL | Following observations of the use of Ozurdex in the clinical practice, the MAH proposed to omit the intravitreal injection procedure video, which was no longer considered necessary to educate treating physicians on the correct application procedure. The CHMP agreed to the removal of the video, and thus the update of Annex II, and considered that the remaining material (consisting of a poster, an injection guide and a copy of the SmPC) was sufficient. In addition, the combination of the educational sets for both indications in uveitis and retinal vein inclusion and streamlining of the information on adverse drug reactions, as proposed by the MAH, was considered by the CHMP to help avoiding the provision of frequent updates and repetitive information to patients and physicians. Other updates to the risk management plan were considered in line with previous requests by the CHMP and PRAC. Minor changes to the package leaflet to better reflect the information in the SmPC and to update the list of local representatives were agreed as well. |

| IA/0014   | A.5.b - Administrative change - Change in the name<br>and/or address of a manufacturer of the finished<br>product, including quality control sites (excluding<br>manufacturer for batch release)  | 10/06/2013 | n/a        |  |  |
|-----------|---|------------|------------|--|--|
| T/0012    | Tansfer of MA from Allergen Pharmaceuticals Ireland<br>(901969) to Allergen Pharmaceuticals Ireland<br>(514125)<br>Transfer of Marketing Authorisation  | 05/02/2013 | 06/03/2013 |  |  |
| IB/0011   | B.II.b.5.a - Change to in-process tests or limits<br>applied during the manufacture of the finished<br>product - Tightening of in-process limits  | 15/01/2013 | n/a        |  |  |
| IB/0010/G | This was an application for a group of variations.<br>B.II.b.3.z - Change in the manufacturing process of<br>the finished product - Other variation<br>B.II.b.5.a - Change to in-process tests or limits<br>applied during the manufacture of the finished<br>product - Tightening of in-process limits | 21/11/2012 | n/a        |  |  |
| IAIN/0009 | C.I.z - Changes (Safety/Efficacy) of Human and<br>Veterinary Medicinal Products - Other variation   | 29/10/2012 | n/a        |  |  |
| IB/0008/G | This was an application for a group of variations.<br>B.II.a.3.b.6 - Changes in the composition<br>(excipients) of the finished product - Other excipients<br>- Replacement of a single excipient with a  | 24/09/2012 | n/a        |  |  |

|         | comparable excipient with the same functional<br>characteristics and at a similar level<br>B.II.e.6.b - Change in any part of the (primary)<br>packaging material not in contact with the finished<br>product formulation - Change that does not affect<br>the product information<br>B.II.e.6.b - Change in any part of the (primary)<br>packaging material not in contact with the finished<br>product formulation - Change that does not affect<br>the product formulation - Change that does not affect<br>the product information<br>B.II.e.3.b - Change in test procedure for the<br>immediate packaging of the finished product - Other<br>changes to a test procedure (including replacement<br>or addition)<br>B.II.e.7.b - Change in supplier of packaging<br>components or devices (when mentioned in the<br>dossier) - Replacement or addition of a supplier |            |            |  |  |
|---------|--|------------|------------|--|--|
| II/0006 | Update of sections 4.3, 4.4 and 4.8 of the SmPC in<br>order to:<br>- add two contraindications (section 4.3)<br>• Aphakic eyes with rupture of the posterior<br>lens capsule<br>• Eyes with Anterior Chamber Intraocular Lens<br>(ACIOL) and rupture of the posterior lens capsule;<br>- amend the wording of section 4.4 following the<br>addition of the new contraindications in section 4.3;<br>- add 2 new ADRs (section 4.8) "Hypotony of eye<br>(associated with vitreous leakage due to injection)"<br>and "Complication of device insertion (implant<br>misplacement)" and amend the description of   | 19/07/2012 | 10/09/2012 | SmPC, Annex<br>II, Labelling<br>and PL | Based on post marketing case reports involving device<br>dislocation the MAH updated the description of the following<br>side effect "device dislocation (device migration)" by<br>amending the text as follows " with or without corneal<br>oedema" in order to better reflect the pathology associated<br>with device migration. In some of the reported cases the<br>eyes were either without lens or were having the natural<br>ocular lens replaced with man-made intraocular lens. The<br>aim of these two contraindications is to minimize the risk of<br>device migration in patients either lacking or having<br>replaced the natural ocular lens. Based on this new safety<br>data the MAH included two new contraindications in<br>patients lacking the crystalline lens or with implanted lens |

|         | "Device dislocation (migration of implant)" ADR by<br>adding "with or without corneal oedema".<br>C.I.3.b - Implementation of change(s) requested<br>following the assessment of an USR, class labelling, a<br>PSUR, RMP, FUM/SO, data submitted under Article<br>45/46, or amendments to reflect a Core SPC -<br>Change(s) with new additional data submitted by the<br>MAH  |            |            | and with rupture of the posterior lens capsule. Based on<br>case reports of complication of device insertion and of<br>decreased intraocular pressure the MAH included in the<br>SmPC the terms 'complication of device insertion (device<br>misplacement)' and "Hypotony of eye (associated with<br>vitreous leakage due to the injection)" respectively as side<br>effects.  |
|---------|---|------------|------------|--|
| IB/0007 | B.II.b.5.a - Change to in-process tests or limits<br>applied during the manufacture of the finished<br>product - Tightening of in-process limits  | 07/09/2012 | n/a        |  |
| II/0005 | The MAH proposed the update of sections 4.4 and<br>4.8 of the SmPC in order to add a warning with<br>regards to implant migration to reflect the changes<br>proposed in PSUR 1 (27 July 2010 to 27 January<br>2011). The MAH also proposed the update of section<br>4.2 of the SmPC in order to include advice on the use<br>of povidone iodine drops 5% to disinfect the<br>periocular skin, eyelid and ocular surface prior to the<br>injection procedure. The Package Leaflet was<br>proposed to be updated in accordance.<br>In addition, the MAH took the opportunity to update<br>the list of local representatives in the Package<br>Leaflet.<br>The requested variation proposed amendments to<br>the SmPC, and Package Leaflet.<br>C.I.3.b - Implementation of change(s) requested | 17/11/2011 | 22/12/2011 | Within this variation the Company has included in the<br>SmPC, sections 4.4 and 4.8, a warning concerning the risk<br>for implant migration from the posterior chamber of the<br>eye into the anterior chamber. The need for such update<br>was identified based on the review of a total of eight<br>medically confirmed cases of device dislocation that have<br>been reported with Ozurdex since the first marketing<br>approval was obtained.<br>The Company, taking advantage of this procedure, has also<br>included in section 4.2 of the SmPC a piece of advice on the<br>use of povidone iodine drops 5% to disinfect the periocular<br>skin, eyelid and ocular surface prior to the injection<br>procedure. The proposed advice regarding the use of<br>povidone-iodine in section 4.2 of the SmPC promotes the<br>safe use of the implant and is in-line with administration in<br>the pivotal clinical trials. |

|           | following the assessment of an USR, class labelling, a<br>PSUR, RMP, FUM/SO, data submitted under Article<br>45/46, or amendments to reflect a Core SPC -<br>Change(s) with new additional data submitted by the<br>MAH   |            |            |                          |  |
|-----------|---|------------|------------|--------------------------|--|
| IA/0004/G | This was an application for a group of variations.<br>B.II.e.6.b - Change in any part of the (primary)<br>packaging material not in contact with the finished<br>product formulation - Change that does not affect<br>the product information<br>B.III.1.a.2 - Submission of a new or updated Ph. Eur.<br>Certificate of Suitability to the relevant Ph. Eur.<br>Monograph - Updated certificate from an already<br>approved manufacturer | 15/08/2011 | n/a        |                          |  |
| II/0001   | Extension of indication to include treatment of adults<br>patients with inflammation of the posterior segment<br>of the eye presenting as non-infectious uveitis.<br>C.I.6.a - Change(s) to therapeutic indication(s) -<br>Addition of a new therapeutic indication or<br>modification of an approved one   | 14/04/2011 | 16/06/2011 | SmPC, Annex<br>II and PL | Please refer to the scientific discussion Ozurdex H-01140-<br>II-001-AR. |
| N/0002    | Minor change in labelling or package leaflet not connected with the SPC (Art. 61.3 Notification)  | 06/04/2011 | n/a        | PL                       |  |