**Medical Device Adverse Event (AE) Report From**

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| AR Report Ref  |
| Report No. (Official Use Only)  |

For use by **Authorised Representatives (AR)** to report events that have take place in:

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| 1. **ADMINISTRATIVE INFORMATION**
 |
| 1. Report Type (select one):[ ]  Initial [ ]  Follow-up [ ]  Final [ ]  Trend  |
| 2. Classification of Event:[ ]  Serious Public Health Concern [ ]  Death [ ]  Serious Injury [ ]  Minor injury[ ]  Other Reportable Event  |
| 3. Date of this report (dd-mmm-yyyy) | dd-mmm-yyyy |
| 4. Date of adverse event (dd-mmm-yyyy) | dd-mmm-yyyy |
| 5. AR awareness date (dd-mmm-yyyy) | dd-mmm-yyyy |
| 6. Expected date of next report (dd-mmm-yyyy) | dd-mmm-yyyy |
| Particulars of the AR Submitting this Report: |
| 7. Name |       |
| 8. Company |       |
| 9. Address |       |
| 10. Mobile Phone No |       |
| 11. Fax |       |
| 12. E-mail |       |
| 13. Other Regulatory Authorities to which this report was *also* sent |
|       |
| **II. CLINICAL EVENT INFORMATION** |
| 1. Event Description: |
| 2. No. of affected people involved |       | 3. No. of devices involved |       |

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| **III. HEALTHCARE FACILITY INFORMATION (OPTIONAL)** |
| 1. Name of the Facility |       |
| 2. Name of Contact Person |       |
| 3. Facility Report No. |       |
| 4. Address |       |
| 5. Phone |       | 6. Fax |       |
| 7. E-mail |       |
| **IV. DEVICE INFORMATION**  |
| Device Information |
| 1. Device Name |       |
| 2. Product License No. |       |
| 3. Product Registration No. |       |
| 4. Nomenclature System | AMDNS / UMDNS Code       |
|  | GMDN Code       |
| 5. Catalogue No. |       |
| 6. Serial No. |       |
| 7. Lot / Batch No. |       |
| Legal Manufacturer Information |
| 8. Name |       |
| 9. Contact Person |       |
| 10. Address |       |
| 11. Phone |       | 12. Fax |       |
| 13. E-mail |       |
| 14. Operator of device at the time of the event[ ]  Healthcare Professional [ ]  Patient [ ]  Other [ ] None |
| 15. Usage of Device[ ]  Initial Use [ ]  Reuse of Single-Use Device [ ]  Reuse of Reusable Devices [ ]  Re-serviced / Refurbished[ ]  Other, please specify: |
| 16. Device Disposition / Current Location:      |

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| **V. RESULT OF MANUAFCTURER’S INVESTIGATION** |
| 1. Manufacturer’s Device Analysis Results: |
| 2. Remedial Action / Corrective Action / Preventive Action: |

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| **VI. INFORMATION OF PATIENT (OPTIONAL)** |
| 1. Age at time of event (months, years) |       |
| 2. Gender (M/F) |  | 3. Weight (kg) |       |
| 4. List of devices involved with the patient (see Section IV):      |
| 5. Corrective action taken relevant to the care of the patient:      |
| 6. Patient outcome:      |
| **VII. OTHER REPORTING INFORMATION (OPTIONAL)** |
| Any events with this device with the same root cause?[ ]  Yes, please specify the rate:       [ ]  No |
| **VIII. COMMENTS**  |
|       |
| **IX. SUBMISSION OF REPORT**  |
| By Mail:      |
| By Fax: (   )      | By e-mail:      |
| **X. DISCLAIMER** |
| Submission of this report does not constitute an admission of manufacturer, AR, user, or patient liability for the event and its consequences. It does not, in itself, represent a conclusion by the AR that the content of this report is complete or confirmed, that the device(s) listed failed in any manner. It is also not a conclusion that the device(s) caused or contributed to the adverse event. |

**GUIDANCE FOR FILLING IN THE ADVERSE EVENT REPORT FORM**

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| **GENERAL** |
| All fields must be completed with appropriate information or “NA” if not applicable to the event or “unknown” when the data is not available. |
| “AR Report No.” on the top right hand corner of the first page is the unique number assigned by the AR to identify the report in the AR’s internal system |
| Reasonable effort must be made to address all elements. However, failure or inability to do so is not justification for failing to submit a report within the establishment timeframes. |
| All GHWP documents referred to in this guidance are available at the GHWP homepage: <http://www.ghwp.info/>  |
| **I. ADMINISTRATIVE INFORMATION** |
| 1. Report Type**Initial:** defined as the first information submitted by the AR about a reportable event, but the information is incomplete and supplementary information will need to be submitted. This includes immediate submission.**Follow-up**: defined as a report that provides supplemental information about a reportable event that was not previously available.**Final:** defined as the last report that the AR expects to submit about the reportable event. A final report may also be the first report.**Trend:** defined as information supplied as a result of significant increase in the rate of (i) reportable events, (ii) non-reportable adverse events, or (iii) adverse events scheduled for periodic reporting. Please refer to the related GHWP guidance document(s). |
| 2. Classification of EventAdverse events that resulted in (i) serious public health concern shall be reported *within 48 hours*, (ii) death, (iii) serious injury shall be reported as soon as possible, but not later than *10 elapsed calendar days* following the awareness of the event.All other reportable events shall be reported as soon as possible, but not later than *30 elapsed calendar days* following the awareness of the event. |
| Please note that the following use errors are reportable events1. Use errors that results in death or serious injury or serious public health concern;
2. When the AR or manufacturer notes a change in trend or a change in pattern of an issue that can potentially lead to death or serious injury or public health concern;
3. When the AR or manufacturer initiates corrective action to prevent death or serious injury or serious public health concern.

Other use errors that do not result in death or serious injury or serious public health concern need not be reported.For details on reportable and non-reportable events, please refer to the related guidance notes. |
| 3 – 6. Dates of this report, date of adverse event, AR awareness date, and expected date of next reportAll dates must be formatted as follows: 2 digit day, 3 letter month, 4 digit year, e.g. 01-JAN-2001Expected date of next report: the date when further information will be provided. This should be “NA” for final report. |
| 7 – 12. Particulars of the AR Submitting this ReportPlease fill in the contact details of the AE’s reporter. |
| 13. Other Regulatory Authorities to which this report was also sentPlease identify to what other regulatory authorities, such as the FDA (US), MHRA (UK), this report was also sent. |
| **II. CLINICAL EVENT INFORMATION** |
| 1. Event DescriptionClarification or relevant information that might impact the understanding or evaluation of the adverse event AND that is not included elsewhere in the report. E.g. “the patient was confused prior to becoming trapped in the bedsides”, “the patient was a very low birth weight premature delivery and had a central line placed three days before onset of cardiac tamponade”, “the X-ray machine was over 20 years old and had been poorly maintained at the time of the adverse event”, etc. |
| 2. No. of affected peopleIncludes any affected individual, e.g. user, patient, or third party |
| 3. No of devicesPlease state the number of devices involved in this event.  |
| **III HEALTHCARE FACILITY INFORMATION** |
| Please provide information about the place of the event. It could include home care, transport or emergency care site. Information in this section is **optional.** |
| **IV. DEVICE INFORMAITON** |
| 1-13. Device information:Please provide information on the device involved. Please repeat this section for each device in separate sheets. |
| 14. Operator of device at the time of the eventPlease indicate the type of operator of the device at the time of the event. “None” means that the problem is noted prior to use. |
| 15. Usage of DevicePlease indicate the usage of the device involved |
| 16. Device Disposition / Current Location:Please provide information on whether and in what state the device is at the time of the report, e.g. “the device has been destroyed”; “the device remains implanted in patient”, “the device was returned to the manufacturer”, the device remains under investigation”, etc. |
| **V. RESULTS OF MANUFACTURER’S INVESTIGATION** |
| 1. Manufacturer’s Device Analysis Results:Specify, for this event, details of investigation methods, results and conclusions.Alternatively, manufacturer’s device analysis report may be submitted. |
| 2. Remedial Action / Corrective Action/ Preventive ActionSpecify if action was taken by manufacturer and/or AR for the reported specific event or for all similar types of products. Include what action was taken by the manufacturer and/or AR to prevent recurrence. Clarify the timeframes for completion of various action plans. |
| **VI. INFORMATION OF PATIENT (OPTIONAL)** |
| Please provide individual patient information (including information of any affected individual, e.g. user, patient, or third party) for each element as appropriate. Please repeat this section for each patient involved in separate sheets.Please note that in some cases, the patient’s age, gender and weight might be irrelevant. In some cases, they are essential, e.g. the age and weight of the patient in regards to some implants.Some events are caused by the combined action of two or more devices, medical or non-medical. Please provide a brief list of devices involved.Information in this section is **optional**. |
| **VII. OTHER REPORTING INFORMATION (OPTIONAL)** |
| If the manufacturer or the AR is aware of similar events with this device with the same root cause, please provide the number of such events. The number should be specified in terms of event per unit sold, or the number of event per unit sold / in use in a region, etc. Information in this section is optional, and is applicable for **final reports** only. |
| **VIII. COMMENTS** |
| Please provide any additional details that are relevant and not requested elsewhere in this report. |