

<b>EudraCT Number:</b> <input type="text"/>	<b>Participant Gender:</b> <input type="text"/>	<b>Participant Date of Birth:</b> <input type="text"/>	<b>Date of sending report to CTO:</b> <input type="text"/>
<b>Participant Randomisation</b> <input type="text"/>	<b>Participant</b> <input type="text"/>	<b>Study Title (short):</b> <input type="text"/>	

1. **What are you reporting:**      SAE       SUSAR\*       Pregnancy       SAR       IME\*\*

*\*Note: If you are reporting a SUSAR the randomisation code for this patient will have to be unblinded.      \*\*Important Medical Event which correlates to eSUSAR E2B criteria "Other Medically Important Condition"*

2. **Report Type:**      Initial Report       Follow up Report       Follow up Report #: \_\_\_\_\_

3. **Protocol Title and Version Number:**

**Evaluation of Event**

4. **Diagnosis (please use medDRA term if known)**

4a. **medDRA Term and code if known:**

5. **Chief or Principal Investigator:**

6. **Sponsor:**

7a. **Date of Onset:**  
(dd/mmm/yy)

7b. **Time of Onset:**  
(if available; hh:mm)

8. **Date person completing report became aware of event:**

(dd/mmm/yy)

9. **Criteria for definition as Serious:**
- Resulted in Death
  - Life threatening
  - In-patient hospitalization or prolongation
  - Persistent or significant disability
  - Congenital anomaly/birth defect
  - N/A (IME or Pregnancy only)

*If there is more than one criterion, choose the more/most significant one. Seriousness is a regulatory definition and should not be confused with severity.*

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10. **Describe Event:** (A summary of signs and symptoms, diagnosis, treatment of event, concurrent treatment, other relevant medical history, including re-challenge details if applicable. Please include the point in the study at which the event occurred.)

11. In the Investigator's opinion was the event related to the Investigational Medicinal Product?

- Definitely\*
- Likely\*
- Possibly\*
- Unlikely
- Not Related

\* This will be reported as a SAR

12. Action Taken With Study Drug

- None
- Dose temporarily reduced
- Dose reduced
- Discontinued temporarily
- Discontinued

13. If related to IMP was this reaction unexpected (Suspected Unexpected Serious Adverse Reaction – SUSAR)

- Yes
- No
- Not Applicable

14. Did event/reaction abate after stopping drug?

- Yes
- No
- Not Applicable

15. Did event/reaction reappear after reintroduction of drug?

- Yes
- No
- Not Applicable

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**16. IMP & Concomitant Medication Information**

<b>Drug Details</b> <i>(include daily dose(s) &amp; generic name)</i>	<b>Therapy Start Date</b> <i>(dd/mmm/yy)</i>	<b>Therapy End Date</b> <i>(dd/mmm/yy)</i>	<b>Date of dose prior to SAE onset</b> <i>(dd/mmm/yy)</i>	<b>Route(s) of administration</b>	<b>Indications for Use</b>
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17. Have Urgent Safety Measures been implemented?

- Yes
- No
- Not Applicable

*If yes, please detail below:*

### Outcome of Event

18. What is the outcome of the SAE?

- Recovered
- Recovered with sequelae
- Continuing
- Resulted in Death
- Unknown

19. Date event resolved: (dd/mmm/yy)

20. Date patient died: (dd/mmm/yy)

21. Cause of death obtained from (if patient died):

- Coroner's inquest
- Death Certificate
- Working diagnosis

### Contact & Signatures

*Please supply contact details where further information may be obtained:*

22. Person to contact:

22a. Centre (if multicentre trial):

23. Phone number:

24. Email address:

\_\_\_\_\_  
Signature (person completing report)

\_\_\_\_\_  
Print Name

\_\_\_\_\_  
Date

\_\_\_\_\_  
Principal Investigator Signature (If multicentre trial)

\_\_\_\_\_  
Print Name

\_\_\_\_\_  
Date

\_\_\_\_\_  
Chief Investigator Signature (If not completing report)

\_\_\_\_\_  
Print Name

\_\_\_\_\_  
Date