



AURORA Serious Adverse Event Report Form

Please email ctu@soton.ac.uk within 24 hours of identification of event FAO: Quality & Regulatory Team
If you do not receive confirmation of receipt within 1 working day from SCTU, please call/re-send the report.

REC No.: 22/LO/0272 Country: UK

A) SITE DETAILS 1. Name of Site <hr style="border-top: 1px dashed black;"/> 2. Name of Principal Investigator	B) WHY WAS THE EVENT SERIOUS? 3. Please select from the list below <input type="checkbox"/> 1 = Resulted in death 2 = Life-threatening 3 = Required inpatient hospitalisation or prolongation of existing hospitalisation 4 = Persistent or significant disability/incapacity 5 = Congenital anomaly/birth defect 6 = Other, Important medical event	C) PATIENT DETAILS 4. Patient Initials <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> 5. Patient ID <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> - <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> 6. Year of Birth <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> 7. Height (cm) <input type="text"/> <input type="text"/> <input type="text"/> 8. Weight (kg) <input type="text"/> <input type="text"/> <input type="text"/> 9. Gender <input type="checkbox"/> Male <input type="checkbox"/> Female Batch Lot Number: _____
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D) DETAILS OF SAE						
10. Date SAE was identified by site <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/>	11. Date SAE sent to SCTU <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/>	13. Grade (CTCAE v5) 1 = Mild 2 = Moderate 3 = Severe 4 = Life-threatening 5 = Death related to AE	14. Date of SAE onset <i>dd- mmm-yyyy</i>	15. SAE status 1 = Resolved 2 = Resolved with sequelae 3 = Ongoing 4 = Worsened 5 = Fatal	16. Date SAE resolved <i>dd- mmm-yyyy</i>	17. Corresponding AE No. As recorded on the AE eCRF
12. Main adverse event diagnosis / symptom - Please record medical term _____ _____ _____		Associated Symptoms (if applicable) - If associated symptoms meet the serious criteria in their own right, a separate SAE will need to be initiated (e.g. if associated symptom is life-threatening or caused prolongation of hospitalisation) _____ _____ _____				

E) TRIAL MEDICATIONS					Investigator's Opinion	
18. Cycle No (delete as appropriate): <input type="text"/>						
19. Trial Treatment name	20. Date of first administration	21. Dose given at most recent administration	22. Date of most recent administration prior to SAE onset	23. Action taken due to SAE 0 = None 1 = Dose reduction 2 = Treatment delayed 3 = Dose reduced & Treatment delayed 4 = Treatment stopped	24. Causal relationship to SAE Is this event causally related to the trial treatment? Definitely <input type="checkbox"/> Probably <input type="checkbox"/> Possibly <input type="checkbox"/> Unlikely <input type="checkbox"/> Unrelated <input type="checkbox"/>	25. PI signature Please provide signature of the PI/delegate who carried out the assessment and date of assessment ----- Signature ----- Date (dd-mmm-yyyy)
Atezolizumab	DD-MMMM-YYYY	1680mg	DD-MMMM-YYYY			

26. Did symptoms abate after stopping drug?	Yes <input type="checkbox"/>	No <input type="checkbox"/>	N/A <input type="checkbox"/>	27. Did symptoms reappear after reintroduction of drug?	Yes <input type="checkbox"/>	No <input type="checkbox"/>	N/A <input type="checkbox"/>
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Patient ID

A	U	2	-							
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Investigator's Opinion

F) OTHER TREATMENTS - do not include therapy given for management of SAE (If none please tick N/A) **N/A**

28. Treatment (Give generic name) - <i>Please include concomitant medication, radiotherapy, surgery and palliative care.</i>	29. Total daily dose	30. Route 1 = Oral 2 = Intravenous 3 = Subcutaneous 4 = Other	31. Start date <i>dd-mmm-yyyy</i>	32. Ongoing 0 = No 1 = Yes	33. End date <i>dd-mmm-yyyy</i>	34. Action taken 0 = None 1 = Dose reduction 2 = Treatment delayed 3 = Dose reduced & treatment delayed 4 = Treatment stopped	35. Causal relationship 1 = Related 2 = Unrelated	36. Sign off Please provide PI (or delegate) signature to confirm they agree with all information listed in section F.
								----- Signature ----- Date (dd-mmm-yyyy)

G) CAUSALITY NOT RELATED TO TREATMENT

37. If you do not consider the main event to be caused by the trial treatment or concomitant medications please **specify any other possible cause** (e.g. medical history, drug or alcohol abuse, family history, findings from special investigations)

N/A

H) DIAGNOSTIC TESTS – If none please tick N/A **N/A**

Test name	Date	Normal range	Results (+ units)



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I) DESCRIPTION OF SAE & RELEVANT MEDICAL HISTORY

Patient ID :

A	U	2	-						
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38. Describe serious adverse event (include manifestation & progression of event and any treatments given in response to the event).

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J) SIGN OFF

Form completed by: _____ Date: _____
Signature *Print Name* *dd-mmm-yyyy*

Contact E-mail: _____ Contact Telephone No: _____

Principal Investigator or delegate: _____ Date: _____
Signature *Print Name* *dd-mmm-yyyy*

Clinical Reviewer Opinion	Definitely, Probably, Possibly – SAR/SUSAR			Unlikely, Unrelated – SAE		SCTU Event No. <table border="1"><tr><td></td><td></td><td></td><td></td></tr></table>				
Assessment against: <i>Please tick the most appropriate status</i>	Definitely	Probably	Possibly	Unlikely	Unrelated					
Atezolizumab										

CR comments:

Signature: _____ Date of Review:

D	D	M	M	M	Y	Y	Y	Y
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