

ADVERSE EVENTS FORM

ID NUMBER: FORM CODE: AES VERSION: 2.0 1/9/2020 Event:
0a) Date of Collection / / / / / / / / Ob) Staff Code / Instructions: This form should be completed if a participant has an adverse event.
Instructions: This form should be completed if a participant has an adverse event.
1) Which study visit is this Adverse Event associated with? Clinic Visit 5 ₁₁ Exacerbation Visit 5 ₁₂ Bronchoscopy Visit 5 ₁₃ Heart Failure Visit ₁₄
2) Adverse Event:
2a) Start Date: /
2b) Stop Date: /
 Severity Mild₁ Event results in mild or transient discomfort, not requiring intervention, or treatment; does not limit or interfere with daily activities (e.g., insomnia, mild headache). Moderate₂ Event is sufficiently discomforting so as to limit or interfere with daily activities; may require interventional treatment (e.g., fever requiring antipyretic medication). Severe₃ Event results in significant symptoms that prevents normal daily activities; may require hospitalization or invasive intervention (e.g., anemia resulting in blood transfusion).
2d) Outcome of Adverse Event Resolved, No Sequelae ₁ Still present-no treatment ₂ Still present-being treated ₃ Residual effects present-not treated ₄ Residual effects present-treated ₅ Death ₆

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2e) Was the ☐ No₀ ☐ Yes		rse Eve	ent expe	cted?			
2f) Was the No ₀ Yes		rse Eve	ent serio	us?			
2g) Please	provid	e a narr	ative de	escription	of the event:		

END OF FORM