AEFI INVESTIGATION FORM

MINISTRY OF HEALTH-GHANA HEALTH SERVICE/FOOD AND DRUGS AUTHORITY

(Only for Serious Adverse Event Following Immunization – Death / Disability / Hospitalization / Cluster)

Section A Basic details						
Region	ion District Case ID					
Place of vaccination (✓): ☐ Govt. health facility ☐ Private health facility ☐ Other (specify) Vaccination in (✓): ☐ Campaign ☐ Routine ☐ Other (specify)						
Address of vaccinati	on site:					
Name of Reporting C	Officer:		Date of investigation Date of filling this f	on: / / orm: /		
•	Designation / Position: This report is: First Interim Final					
Telephone # landline Patient Name:	(with code).	Mobi	ile.	e-mail:	Sex: M F	
(use a separate form for ea	ch case in a cluster)					
Date of birth (DD/MM/	YYYY): /	_ /				
Years >18-60 Ye	_ years months _ ears >60 Years				ears	
Patient's full address v	with landmarks (Street	name, house numbe	er, locality, phone nur	nber etc.):		
Name of vaccines/diluent received by patient	Date of vaccination	Time of vaccination	Dose (e.g. 1 st , 2 nd , etc.)	Batch/Lot number	Expiry date	
- 1				Vaccine Diluent	Vaccine Diluent	
				Vaccine Diluent	Vaccine Diluent	
				Vaccine	Vaccine	
				Diluent Vaccine	Diluent Vaccine	
				Diluent	Diluent	
				Vaccine Diluent	Vaccine Diluent	
Type of site (✓) ☐ Fixed ☐ Mobile ☐ Outreach ☐ Other Date of first/key symptom (DD/MM/YYYY): / Time of first symptom (hh/mm): / Date of hospitalization (DD/MM/YYYY): / Date first reported to the health authority (DD/MM/YYYY): / /						
Status on the date of investigation (✓): ☐ Died ☐ Disabled ☐ Recovering ☐ Recovered completely ☐ Unknown						
If died, date and time of death (DD/MM/YYYY): / / (hh/mm): / Autopsy done? (✔) ☐ Yes (date) ☐ No ☐ Planned on (date) TimeAttach report (if available)						
Section B Relevant patient information prior to immunization						
Doot history of the "	Criteria		Finding		f yes provide details)	
Past history of similar		1	Yes / No / Unkn Yes / No / Unkn			
Adverse event after previous vaccination(s) History of allergy to vaccine, drug or food			Yes / No / Unkn			
Pre-existing comorbidity/ congenital disorder?			Yes / No / Unkn			
		Yes / No / Unkn				
Pre-existing acute illness (30 days) prior to vaccination? Has the patient tested Covid19 positive prior to vaccination? Yes / No / Unkn Yes / No / Unkn						
Pre-existing illness (30 days) / congenital disorder Yes / No / Unkn						
	ntion in last 30 days, w		Yes / No / Unkn			
Was the patient receiving any concomitant medication? Yes / No / Unkn						
(If yes, name the drug, indication, doses & treatment dates)						
Patient currently on concomitant medication? (If yes, name the drug, indication, doses & treatment dates) Family history of any disease (relevant to AEFI) or allergy Yes / No / Unkn						
Family history of any For adult women	uisease (relevant to F	LETI) OF Allergy	Yes / No / Unkn			
	egnant? Yes (weeks)	Currently pregnant? Yes (weeks) / No / Unknown				

Name	Case ID Number	AEFI Investigation Page 2/5					
Currently breastfeeding? Yes / No							
For infants The birth was ☐ full-term ☐ pre-term ☐ post-term. Birth weight:							
Delivery procedure was ☐ Normal ☐ Caesare	an Assisted (forceps, vacuum etc	:.)					
Section C Details of first 6	examination** of serious AEF	l case					
Source of information (✓ all that apply): ☐ Examination ☐ Other If from	on by the investigator	ents					
Name of the person who first examined/treated the patient: Name of other persons treating the patient:							
Other sources who provided information (specify):							
Signs and symptoms in chronological order from the	time of vaccination:						
Name and contact information of person completing these clinical details:	Designation:	Date/time					
summary, laboratory reports and autopsy reports attached documents below • If patient has not received medical care – obta additional sheets if necessary)							

Name					Investig	nvestigation Page 3/5					
Provisional / Final diagnosis:											
Section D	Detai	ls of vaco	ines pro	vided at t	he site link	ced to A	EFI on tl	ne corres	pondir	ng day	
Number immunized for each antigen at	Vaccine name										
session site. Attach record if available.	Number of doses										
a) When was	the patien	t immunize	d? (√	the 🗌 bel	ow and resp	ond to AL	.L questio	ns)			
☐ Within th	ne first vac	cinations o	f the sessi	on 🗌 With	in the last va	accination	s of the s	ession 🗌	Unknow	n	
In case of I					within the fir	st few do	ses of the	e vial adm	nistered	? 🗌 wi	ithin the
					o recommen	dations fo	or use of t	his		Yes* / N	lo
c) Based on y been unste	rile?				cine (ingredie				e Yes*	Yes* / No / Unable to assess	
					cine's physic e time of adr			olour,	Yes*	Yes* / No / Unable to assess	
 turbidity, foreign substances etc.) was abnormal at the time of administration? e) Based on your investigation, do you feel that there was an error in vaccine reconstitution/preparation by the vaccinator (e.g. wrong product, wrong diluent, improper mixing, improper syringe filling etc.)? 					Yes*	Yes* / No / Unable to assess					
f) Based on your investigation, do you feel that there was an error in vaccine handling (e.g. break in cold chain during transport, storage and/or immunization session etc.)?					Yes*	Yes* / No / Unable to assess					
g) Based on your investigation, do you feel that the vaccine was administered incorrectly (e.g. wrong dose, site or route of administration, wrong needle size, not following good injection practice etc.)?					Yes*	Yes* / No / Unable to assess					
h) Number im											
i) Number im	i) Number immunized with the concerned vaccine in the same session										
	j) Number immunized with the concerned vaccine having the same batch number in other locations. Specify locations:										
k) Is this case a part of a cluster?						Ye	Yes* / No / Unkn				
i. If yes, how many other cases have been detected in the cluster?											
a. Did all the cases in the cluster receive vaccine from the same vial?						Ye	Yes* / No / Unkn				
b.If no, number of vials used in the cluster (enter details separately)											
*It is compulso										_	
Section E I		-	_		ce(s) whe				was us	<u>sed</u>	
(Complete this section by asking and/or observing practice) Syringes and needles used:											
Are AD syringe			ion?							Yes / No	o / Unkn
If no, specify the type of syringes used: ☐ Glass ☐ Disposable ☐ Recycled disposable ☐ Other											
Specific key findings/additional observations and comments:											
Reconstitution: (complete only if applicable, ✓ NA if not applicable)											
Reconstitution. (c)			ioabie, *	14A II 110C	applicable)				St	atus	
Same reconstitution syringe used for multiple vials of same vaccine?					Yes	No	NA				
Same reconstitution syringe used for reconstituting different vaccines? Yes No NA							NΑ				

Name			gation Pa	ige 4/5				
	Separate reconstitution syringe for each vaccine vial?	Yes	No	NA				
	Separate reconstitution syringe for each vaccination?	Yes	No No	NA NA				
Are the vaccines and diluents used the same as those recommended by the manufacturer? Yes								
Specific key findings/additional observations and comments:								
Section F	Cold chain and transport							
	(Complete this section by asking and/or observing practice)							
Last vaccine s								
Is the temper	erature of the vaccine storage refrigerator monitored?		Yes / No					
o If "y	res", was there any deviation outside of 2-8° C after the vaccine was placed inside?		Yes / N	lo				
o If "y	res", provide details of monitoring separately.							
Was the co	rrect procedure for storing vaccines, diluents and syringes followed?		Yes / No / Unkn					
 Was any ot 	her item (other than EPI vaccines and diluents) in the refrigerator or freezer?		Yes / No / Unkn					
 Were any p 	artially used reconstituted vaccines in the refrigerator?		Yes / No	/ Unkn				
Were any u	nusable vaccines (expired, no label, VVM at stages 3 or 4, frozen) in the refrigerator	·?	Yes / No	/ Unkn				
Were any u	nusable diluents (expired, manufacturer not matched, cracked, dirty ampoule) in the	store?	Yes / No	/ Unkn				
	dings/additional observations and comments:							
Vaccine transp	ortation:							
	cine carrier used		Yes / No					
Was the vaccine carrier sent to the site on the same day as vaccination?								
Was the vaccine carrier returned from the site on the same day as vaccination?				Yes / No / Unkn Yes / No / Unkn				
Was a conditioned ice-pack used?								
Specific key find	dings/additional observations and comments:							
Section G	Community investigation (Please visit locality and interview pa	rents/oth	ers)					
Yes / No / Unkn	ar events reported within a time period similar to when the adverse event occurred own If yes, describe:	and in the	e same k	ocality?				
If yes, how man	y events/episodes?							
Vaccinated:Not vaccina	ted:							
Saler deliminant								

Section H

Other findings/observations/comments

Name	Case ID Number	AEFI Investigation Page 5/5