

Detailed Information on (Product) Failures or (Near) Events

Customer Data	Product Data		
Company*	Quantity*		
Salutation Titel	Part number*		
First name* Last name*	Description*		
Street/number*	Serial/Lot number*		
ZIP* City*	Invoice number*		
Phone number*	Purchase date*		
You have rated a (product) failure or a (near) every Please describe the failure or the event in greater			
A. General Data Date of complaint When and how (kind of communication used) was the per	son informed that reported the complaint?		
When and now (kind of communication asca) was the per	soft informed that reported the complaint:		
Name, function and address of reporting person:			
First name Last name	Function		
Street/number	ZIP City		
Was there a formal notification to the competent authority	? Yes No No		

*mandatory field Seite 1 von 4

B. Patient/User Data

If a patient or us	er was concerned	by the reported e	event, we a	sk for the following	anonymized	I data:
Age	Gender	Weight		Special physiologi	cal condition	S
Participation in a	a clinical trial					
More details						
C. Details al	bout the Even	t				
	escriptions of circu					
What kind of am	nbient conditions v	vas the product/m	nodical davi	oo boing used in?		
in closed ro					in an incubat	
		in oper			in an incubat	Or
	escent tube or othe		rce in the ir	nmediate vicinity		
	se provide a pictur					
How was the En	nviteC device or se	nsor involved in th	ne event?			
Country in which	n the event occurre	ed				
If possible, pleas	se attach relevant i	est results and ar	nonymized	patient data to this	report.	
	lical devices involv			•		Yes No
	dicate the following					
Туре	Mo	odel		Serial number		More details
When measuring	g vital parameters:	Which type of mo	onitor was i	used? Please state	the following	g details:
Туре	Mo	odel		Software version		Serial/Lot number
Which failure me	essage was indicat	ed on the monito	r?			

If possible, please provide a p	picture of the monitor with the se	ensor used.	
Has any adapter or extension If yes, please state the following	Yes No		
Туре			
More details			
Who used/applied the medical	al device?		
Job title	Position	Department	
For SpO ₂ sensors: How and h	now often was the sensor repos	sitioned?	
For SpO ₂ sensors: Did the nu	rses use the sensor according t	to EnviteC user instructions?	Yes No
During measurement, were other vital parameters measured at the same time on the same monitor?			Yes No
If yes, which other parameters	s?		
Please indicate the contact pe	erson at the final customer/end	user with contact details:	
First name	Last name	Phone number	E-Mail
Supplemental information			
D. Details about the M	Indical Daviso		
	de the following product details:		
Product description	Min. or maximum shelf life	Serial/Lot number	Part number
] [
Other descriptive details			

The medical product is a:	Disposable product	Reusable product	
	New product	Reprocessed produ	ct
Is the medical device availab	le for analyses?		Yes No
Has the product already been	Yes No		
Supplemental information			
E. Miscellaneous			
If available, please provide co	opies of communication from/to	final customer, end user, op	perator and/or authorities.
Supplemental information			

Please forward the completed document with any attachment to EnviteC-Service@honeywell.com

Your support is highly appreciated!