FSN Ref: 2022-04

Date: 2022-09-01

Urgent Field Safety Notice QTYPE

For Attention of: Users of product QTYPE lots E049, E050, E051, E052, E053, E054, E055, E056, E057, E058, E059 and E060

Contact details (name, e-mail, telephone, address etc.)

Anna Bereza-Jarocinska regulatory-se@caredx.com +46 8 508 939 00 Franzéngatan 5 112 51 Stockholm Sweden

	A. Information on Affordad Doubook				
	1. Information on Affected Devices*				
1.	1. Device Type(s)				
	Olerup QTYPE 11 kits consist of qPCR plates containing pre-aliquoted and dried reaction mixes in each well, together with Master Mix provided in separate vials				
1.	2. Commercial name(s)				
	Olerup QTYPE 11				
1.	3. Unique Device Identifier(s) (UDI-DI)				
	N/A				
1.	4. Primary clinical ouroose of device(s)				
	Olerup QTYPE 11 HLA Typing Kits are qualitative in vitro diagnostic tests for the DNA typing of HLA Class I and Class II alleles. The kits are to be used as an aid in determining HLA-A, 8, C, DR81, DR83, DR84, DR85, DQA1, DQ81, DPA1 and/orDP81 alleles with low to intermediate resolution in human genomic DNA samples extracted from anticoagulated blood, to aid in transfusion and transplantation donor and recipient matching. Olerup QTYPE 11 kits are for professional use only and must not be used as the sole basis for makinQ clinical decisions.				
1.	Device Model/Catalogue/oart number(s)				
	201.701-03/10				
1.	6. Software version				
	N/A				
1.	7. Affected serial or lot number range				
	E049, E050,E051, E052, E053, E054, E055, E056, E057, E058,E059andE060				
1.	Associated devices				
	N/A				

	2. Reason for Field Safety Corrective Action (FSCA)			
2.	Description of the product oroblem			
	A known 8*50:01:01:02 sample was typed as 8*50:02 due to an incorrect reactivity in the mix in H7 FAM. 8*50:01:01:02, 8*50:01:01:08, 8*50:01:08, 8*50:01:18, 8*50:51, and 8*50:57 were not included in the string of potential result options due to the error, which has now been corrected. This applies to all active QTYPE lots (E049-E060).			
2.	Hazard giving rise to the FSCA			
	A known 8*50:01:01:02 sample was typed as 8*50:02 due to an incorrect reactivity in the mix in H7 FAM. The serological split within the 8*50 group caused a serological mistyping.			
2.	2. 3. Probability of problem arisinçi			
	The problem is only seen in samples with the rare alleles 8*50:01:01:02, 8*50:01:01:08, 8*50:01:08, 8*50:01:18, 8*50:51, or 8*50:57.			



FSN Ref: 2022-04

2.	Predicted risk to patient/users	
	Low	
2.	5. Further information to help characterise the problem	
	N/A	
2.	6. Backoround on Issue	
	HLA-8 resuft is reported as 8*50:02 instead of 8*50:01:01:02 due to incorrect reactivity in well H7	
	FAM. The serological split within the 8*50 group caused a serological mistypina.	
2.	7. Other information relevant to FSCA	
	This issue will affect all lots from E049 when analysed with a kit file older than	
	Typinçikit QTYPE 20220825.	

	3. Type of Action to mitigate the risk			
3.	1. Action To Be Taken by the User*			
	Identify Device ☐ Quarantine Device ☐ Return Device ☐ Destroy Device ☐ On-site device modification/inspection ☐ Follow patient management recommendations			
	☐ Take note of amendment/reinforcement of Instructions For Use (IFU) Other ☐ None			
	Describe: Update to typingkitQTYPE_20220825.vda file. Return signed Customer/Distributor Reply Form			
3.	2. By when should the action be completed? Typing kit file to be updated as soon as possible. Completed Customer Reply to be returned by 2022-Sep-16			
3.	Particular considerations for: IVO			
	No			
3.	4. Is customer Reply Required? Yes (If yes, form attached specifyino deadline for return)			

CareDx ·

FSN Ref: 2022-04

3.	5. Action Being Taken by the Manufacturer	
	☐ Product Removal ☐ On-site device modification/inspection	
	☐ Software upgrade ☐ IFU or labelling change	
	IZI Other ☐ None	
	The Olerup QTYPE 11 kit file has been updated and reaction H? FAM has received a specificity chançie in typinçi kit file Typingkit QTYPE 20220825.vda	
3	6. By when should the 2022-Aug-26 action be completed?	
3.	7. Is the FSN required to be communicated to the patient No /lay user?	
3.	8. If yes, has manufacturer provided additional information suitable for the patienUlay user in a oatienUlay or non-professional user information letter/sheet?	
	N/A	

	4. G	eneral Information
4.	1. FSN Type	New
4.	For updated FSN, reference number and date of previous FSN	N/A
4	3. Further advice or information already expected in follow-up FSN?	No
4.	4. Manufacturer information (For contact details refer to paçie 1 of this FSNJ	
	a. Comoanv Name	CareDx AB
	b. Address	Franzénaatan 5, 112 51 Stockholm, Sweden
	c. Website address	www.caredx.com
4.	The Competent (Regulatory) Authority of your country has been informed about this communication to customers.	
4.	6. List of attachments/appendices:	Distributor or Customer Reoly Form
4.	7. Name/Signature	

Transmission of this Field Safety Notice
This notice needs to be passed on all those who need to be aware within your organisation or to any organisation where the potentially affected devices have been transferred. (As appropriate)
Please transfer this notice to other organisations on which this action has an impact. (As appropriate)
Please maintain awareness on this notice and resulting action for an appropriate period to ensure effectiveness of the corrective action.
Please report all device-related incidents to the manufacturer, distributor or local representative, and the national Competent Authority if appropriate, as this provides important feedback.