

FIELD SAFETY NOTICE
ACTION REQUIRED

Rheumatoid Factors 2 (RF 2) test over recovery

May xx, 2020

CUSTOMER INFORMATION

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Dear Valued Customer:

The purpose of this letter is to advise you that Thermo Fisher Scientific Oy, part of Thermo Fisher Scientific Inc., is conducting a Field Safety Corrective Action (FSCA) for the in vitro diagnostic products as listed below (Table 1). Our records indicate that you have purchased units of the affected products.

Table 1. PRODUCT INFORMATION

Product Name	Product code	Lot No.
Rheumatoid Factors 2 (RF 2)	981920	R113 (Exp. 2020-08-31)
Rheumatoid Factors 2 (RF 2)	981920	R315 (Exp. 2020-08-31)
Rheumatoid Factors 2 (RF 2)	981920	Following already expired lots: MA12 (Exp. 2018-04-30) MC28 (Exp. 2018-04-30) N501 (Exp. 2018-04-30) N771 (Exp. 2018-11-30) N949 (Exp. 2018-11-30) NA90 (Exp. 2018-11-30) P132 (Exp. 2018-11-30) P421 (Exp. 2019-12-31) P932 (Exp. 2019-12-31) P932B (Exp. 2019-12-31)
Rheumatoid Factors Control	981252	S376A, S376B (Exp. 2021-05-31)
Rheumatoid Factors Control	981252	Following already expired lots: MA93A (Exp. 2018-04-30) N149A, N149B, N149C (Exp. 2018-11-30) NC04A, NC04B (Exp. 2019-07-30) PA09A, PA09B (Exp. 2020-01-31)

REASON FOR FIELD CORRECTION:

It has been identified that the RF values for the above-listed lots of Rheumatoid Factors 2 (RF2) test (Product Code 981920) calibrator and associated control, Rheumatoid Factors Control (Product Code 981252), have been incorrectly assigned, resulting in over recovery of patient test results to the RF reference material 1st British standard from NIBSC Ref 64/002.

Through this communication, customers are being directed to stop using the impacted RF2 lots and begin using replacement material (Product Code 981920 Lot S642B or later), which has been produced to align more closely with the RF reference material, NIBSC Ref 64/002. Customers can continue using Rheumatoid Factors Control (Product code 981252) Lot S376 with newly reassigned control values to be in line with the adjusted test level as detailed below.

When first testing with the newly released lot (981920 Rheumatoid Factors 2 Lot S642B) or subsequent lots you may observe a decrease of approximately 23 % in patient results and controls when compared to the most recent lots R113 and R315 of the same product.

IMPACT ON PATIENT RESULTS:

With the use of the lots listed in Table 1, there is a risk of reporting incorrect, falsely elevated RF levels, which may lead to unnecessary additional serological testing.

ACTIONS TO BE TAKEN BY THE CUSTOMER / USER:

1. If you have inventory of the above mentioned (Table 1) Thermo Fisher Scientific products, stop using listed lots of the Rheumatoid Factors (RF2) test (Product Code 981920). You can continue using Rheumatoid Factors Control (Product Code 981252) Lot S376 with newly reassigned control values as detailed below.
2. Retain a copy of this letter for your laboratory records.
3. Please contact your local Thermo Fisher Scientific representative for further information and the number of RF2 kits to be credited (kits that you still have in your inventory) by replacement product.
4. As appropriate, contact your Medical Professional for evaluation of further action.
5. If you have any inventory, please destroy affected RF2 products according to local waste management rules.
6. Please, fill out the MEDICAL DEVICE FIELD CORRECTION - Response Form and return it within 5 days of the date of this letter to your distributor as instructed in the form and as listed below:
FAX: [add distributor contact information] or
Email: [add distributor contact information]

TYPE OF ACTIONS TO BE TAKEN BY THE MANUFACTURER:

The calibrator value for product code 981920 (from Lot S642B, Exp. 2021-04-30) and control value for product code 981252 (from Lot S376, Exp. 2021-05-31) have been adjusted to the manufacturer's original test level to realign more closely to the reference material 1st British standard from NIBSC Ref = 64/002.

NOTE: Updated Rheumatoid Factors Control 981252 Lot S376 value sheet is available in e-labeling. Please download your copy from the e-documents using the link and key code given below.

<http://www.e-labeling.eu/TSF>, key code TSF981252VS_S376

We appreciate your immediate attention to this Field Safety Corrective Action. We apologize for any inconvenience this may have caused and appreciate your understanding as we take action to ensure customer safety and satisfaction.

If you have any further questions, please contact your local Thermo Fisher Scientific representative by sending an email to [[add distributor contact information](#)].

Sincerely,

Silja Halme
Director, Quality Assurance & Regulatory Affairs
Thermo Fisher Scientific Oy
Analyzers & Automation
Clinical Diagnostics

**MEDICAL DEVICE FIELD CORRECTION
Response Form**

Rheumatoid Factors 2 (RF 2) test over recovery

I have read and understand the attached Field Safety Notice and field action instructions:
_____ (initials)

I understand that this applies to all inventory of the affected in vitro diagnostic medical device products listed in Table 1 that I have received, and I have destroyed the existing affected RF2 kit inventory (if any): _____ (initials)

Do you have any knowledge of adverse medical events associated with the products listed in this Field Safety Notice?
_____ Yes _____ No

If yes, please explain:

RETURN RESPONSE (please provide additional information, if applicable):

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PLEASE RETURN COMPLETED AND SIGNED FORM TO EMAIL: [\[add distributor contact information\]](#)

Signature of Receipt by Customer: _____

Customer Name/Title:	
Date:	
Company/Institute:	
Telephone:	
Email Address:	

It is important that your organisation takes action as detailed in this letter and also replies without delay by using this response form. Your reply is evidence, which Thermo Fisher Scientific and Regulatory Agencies need to monitor the progress of

FSCAs. Without your reply Thermo Fisher Scientific Oy cannot verify the effectiveness or completeness of this Field Safety Corrective Action.