



Geneva, 4th November 2021
FSCA 5333-1

Please distribute to the Laboratory Manager
– IMPORTANT, GLOBAL FIELD SAFETY NOTICE - Correction
VIDAS® Immuno-Assays Multiple references
Substrate error - Potential delayed results without medical impact –
Response required

Dear bioMérieux Customer,

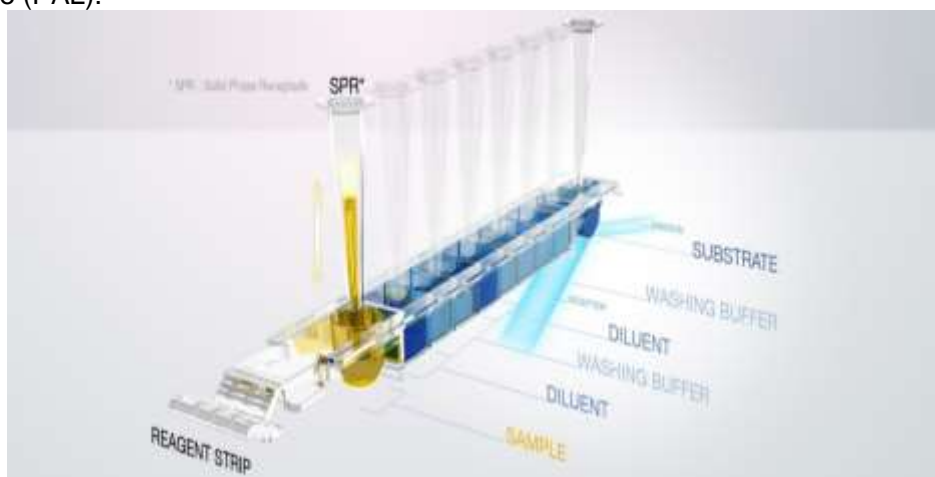
This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

The intent of this letter is to share with you important information regarding clinical VIDAS® Immuno-Assays references products listed below in Table 1. Your laboratory received one/several of the listed clinical VIDAS® Immuno-Assays references products and lots.

Description of the issue

Since July 2021, bioMérieux has been receiving an increasing number of complaints linked to a VIDAS® “substrate error”. It prevents the test from being run, therefore leads to a potential delayed results as you need to run another test.

A measurement of the background noise signal (RFU) is made by the VIDAS® system prior to launching the reaction. An acceptable limit is defined during product design for each reference of finished goods. Three values exist as acceptable limits depending on the assay: 300, 350 and 500 RFU. The substrate is present in the last well of the strip of all VIDAS® immuno-assays and allows fluorescence when degraded by the enzyme (PAL).



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The level of fluorescence is then correlated with the results of all tests.

When you are performing a test, if the RFU is higher than the acceptable limit defined during the product development, there is an error message displayed by the system: "Substrate Error". The test is stopped and this alarm prevents the system to provide any result .

This alarm being present on all the systems of the VIDAS® family, it guarantees that no false results can be given in case of a substrate degradation. This means that there is only a potential risk of delayed results.

Investigations were immediately initiated to identify the root-cause, the following were identified:

- All lots impacted of VIDAS® Immuno-assays were conform to the specifications at release.
- The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using a common lot of raw material (4-MUP) that was identified as the most probable common root-cause.
- The scope of the issue was identified on all lots of VIDAS® Immuno-Assays manufactured using substrate batches containing this concerned lot of raw material. Most of the lots of VIDAS® Immuno-Assays since February 2021 were manufactured using this impacted raw material but not all of them.
- The problem is due to an accelerated degradation of the substrate. It follows a linear model over time leading to RFU acceptable limits being reached before the end of the registered shelf life of the product. This reinforced the reason why the VIDAS® Immuno-Assays with the lowest RFU limit (set at 300), were the first assays impacted : VIDAS® HIV DUO QUICK (Ref. 30447).
- Kinetic evolution analyzes were performed by measuring substrate RFU of a statistically representative number of VIDAS® immuno- assays retained batches (manufactured with the substrate containing the concerned raw material) at different shelf-lives. The model was validated on numerous data (~ 450 000) collected from the field (customers) .
- The analysis of the kinetic model allows us to predict the degradation trend of the substrate using the concerned batch of 4MUP and therefore to revise the expiry dates for each lot of impacted VIDAS® Immuno- assays finished products.
- When used until the revised expiry date, the product continues to perform per its registered performance specifications.
- Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assays products are required to ensure that the specified products will continue to perform per labelled performance specifications.
- Indeed, even if there is no medical impact in case of delayed result when using the impacted lots of clinical VIDAS® Immuno-assays listed in Table 1 below, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

We understand this complex matter requires an extra time commitment from you and your team. We thank you for your cooperation which is essential to ensure the successful implementation of this corrective action, in order to protect the safety of our patients.

To clarify the actions you are required to take, we have provided an attachment for each VIDAS® Immuno-assays product (Table 1) below. The attachments will clearly explain the following information:

- A list of all impacted lot numbers for each clinical VIDAS® Immuno-assay and associated product reference number,
- The revised expiration date for each impacted lot,
- Identification of lots that should be discarded due to revised expiry date,



- Additional actions required to be implemented within your institution.

Please determine which product references you currently have in stock that are referenced in Table 1, and implement the actions defined in the applicable attachments.

We are currently reworking some lots of VIDAS® Immuno-Assays in stock applying a sticker with the revised expiry date on top of the kits. However, to ensure service continuity you may receive, for a short period of time, impacted lots of clinical VIDAS® Immuno-Assays without sticker. All those lots are in the scope of this Urgent Field Safety Notice, and detailed in the different attachments of the table 1.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact the Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely,

bioMérieux (Suisse) SA

A handwritten signature in blue ink that reads "Sabrina Wulf".

Sabrina Wulf
Product Manager Immunoassays

Attachment A: Acknowledgement Form

Table 1: List of impacted clinical VIDAS® Immuno-Assay references (for which a delayed result has no medical impact)

| Product Ref. | Product name | Attachments |
|--------------|-----------------------------------|-------------------|
| 30115 | VIDAS® PROTEIN C 30 TESTS | See attachment 1 |
| 30218 | VIDAS® MUMPS IGG 60 TESTS | See attachment 2 |
| 30219 | VIDAS® MEASLES IGG 60 TESTS | See attachment 3 |
| 30221 | VIDAS® RUB IGG II 60 TESTS | See attachment 4 |
| 30222 | VIDAS® TOXO IGG AVIDITY 30TESTS | See attachment 5 |
| 30226 | VIDAS® RUB IGG 60 TESTS | See attachment 6 |
| 30235 | VIDAS® EBNA IGG 30 TESTS | See attachment 7 |
| 30236 | VIDAS® EBV VCA.EA IGG 30 TESTS | See attachment 8 |
| 30237 | VIDAS® EBV VCA IGM 30 TESTS | See attachment 9 |
| 30305 | VIDAS® HBE.ANTI-HBE 30 TESTS | See attachment 10 |
| 30312 | VIDAS® ANTI-HAV TOTAL 30 TESTS | See attachment 11 |
| 30319 | VIDAS® LYME IGM 60 TESTS | See attachment 12 |
| 30320 | VIDAS® LYME IGG 60 TESTS | See attachment 13 |
| 30400 | VIDAS® TSH 60 TESTS | See attachment 14 |
| 30402 | VIDAS® FT3 60 TESTS | See attachment 15 |
| 30403 | VIDAS® T3 60 TESTS | See attachment 16 |
| 30404 | VIDAS® T4 60 TESTS | See attachment 17 |
| 30406 | VIDAS® LH 60 TESTS | See attachment 18 |
| 30407 | VIDAS® FSH 60 TESTS | See attachment 19 |
| 30410 | VIDAS® PROLACTINE 60 TESTS | See attachment 20 |
| 30411 | VIDAS® FERRITINE 60 TESTS | See attachment 21 |
| 30413 | VIDAS® AFP 60 TESTS | See attachment 22 |
| 30419 | VIDAS® IGE 60 TESTS | See attachment 23 |
| 30420 | VIDAS® B2 MICROGLOBULI 30 TESTS | See attachment 24 |
| 30426 | VIDAS® CA 125II 30 TESTS | See attachment 25 |
| 30427 | VIDAS® CA 19-9 30 TESTS | See attachment 26 |
| 30428 | VIDAS® TPSA 60 TESTS | See attachment 27 |
| 30429 | VIDAS® CA 15-3 30 TESTS | See attachment 28 |
| 30431 | VIDAS® ESTRADIOL II 60 TESTS | See attachment 29 |
| 30436 | VIDAS® VWF 30 TESTS | See attachment 30 |
| 30440 | VIDAS® FPSA 30 TESTS | See attachment 31 |
| 30441 | VIDAS® TSH3 60 TESTS | See attachment 32 |
| 30453 | VIDAS® CEA (S) 60 TESTS | See attachment 33 |
| 30459 | VIDAS® FT4N 60 TESTS | See attachment 34 |
| 30461 | VIDAS® ANTI-TPO 30 T | See attachment 35 |
| 30462 | VIDAS® ANTI-TG 30 T | See attachment 36 |
| 30463 | VIDAS® 25-OH VITAMINE D TOTAL 60T | See attachment 37 |
| 414320 | VIDAS® TESTOSTERONE II 30 TESTS | See attachment 38 |
| 416436 | VIDAS® LYME IGM II 60 TESTS | See attachment 39 |
| 417011 | VIDAS® AMH 30 TESTS | See attachment 40 |



| | | |
|--------|---|-------------------|
| 417401 | VIDAS [®] LYME IGG II 60 TESTS | See attachment 41 |
| 418116 | VIDAS [®] HEV IGG 30T | See attachment 42 |
| 422010 | VIDAS [®] PTH (1-84) 30T | See attachment 43 |
| 423079 | VIDAS [®] ANTI-DENGUE IGG 60 TESTS | See attachment 44 |
| 423111 | VIDAS [®] TB-IGRA 60 TESTS | See attachment 45 |
| 423833 | VIDAS [®] SARS-COV-2 IgM (9COM) 60 T | See attachment 46 |
| 423834 | VIDAS [®] SARS-COV-2 IgG (9COG) 60 T | See attachment 47 |
| 424114 | VIDAS [®] SARS-COV-2 IgG II | See attachment 48 |



**Attachment A: ACKNOWLEDGEMENT FORM
URGENT: FIELD SAFETY NOTICE**

**FSCA 5333-1 - VIDAS® Immuno-Assays Multiple references
Substrate error - Potential delayed results without medical impact –
Response required**

In compliance with legal traceability requirements we thank you for completing this form even if you no longer have this reagent

**Please send us back the completed acknowledgment of receipt by email
(ch_support@biomerieux.com).**

Name of the laboratory: _____

Contact person: _____

City and postal Code: _____

Customer No.: _____

For products with No remaining shelf life, we ask you to enter the number of destroyed kits in the following table.

| REF | Lot # | Product Name | Number of kits destroyed |
|-----|-------|--------------|--------------------------|
| | | | |
| | | | |
| | | | |
| | | | |
| | | | |

With your signature you confirm that you have received the Urgent Field Safety Notice „ **VIDAS® Immuno-Assays Multiple references – Substrate error - Potential delayed results without medical impact –** ”.

You further confirm that you will follow the instructions and implement the actions as indicated in the Urgent Field Safety Notice.

Have you encountered impact on patients' results, or reports of illness or injury related to the identified issue ?

No

Yes

If **YES**, please give your telephone number for contacting you: _____

DATE

SIGNATURE



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 1 related to FSCA 5333-1- VIDAS® PROTEIN C (Ref. 30115) - Substrate error Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1 and Table 2, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1 and 2, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1 and 2, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1 or Table 2.

- **For products with No remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify impacted lots in Table 1,
 - Stop using the listed impacted lots
 - Destroy the remaining lots in your inventory.
- **For products with remaining shelf life (taking into account the revised expiry date) in Table 2:**
 - Identify products listed in Table 2,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 2.
 - Continue product use until revised expiry date.
- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® PROTEIN C 30 TESTS (Ref. 30115) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible. We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
 bioMérieux (Suisse) SA

Table 1: Product with NO remaining shelf life (taking into account the revised expiry date).

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|--------------|---------------------------|---------------------|---------------------|
| 30115 | 1008662180** | VIDAS® PROTEIN C 30 TESTS | 19-Mar-22 | 18-Oct-21 |
| 30115 | 1008571770** | VIDAS® PROTEIN C 30 TESTS | 5-Feb-22 | 6-Sep-21 |

** Lots under PSS (Product Stop Shipment)

Table 2 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|---------------------------|---------------------|---------------------|
| 30115 | 1008801740 | VIDAS® PROTEIN C 30 TESTS | 26-May-22 | 12-Dec-21 |
| 30115 | 1008881620 | VIDAS® PROTEIN C 30 TESTS | 19-Jul-22 | 12-Feb-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 2 related to FSCA 5333-1- VIDAS® MUMPS IGG 60 TESTS (Ref. 30218) - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.
- Contact your local bioMérieux representative to order the replacement products when appropriate,



- As a reminder, please store the VIDAS® MUMPS IGG 60 TESTS (Ref. 30218) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|---------------------------|---------------------|---------------------|
| 30218 | 1008769120 | VIDAS® MUMPS IGG 60 TESTS | 22-Apr-22 | 19-Apr-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 3 related to FSCA 5333-1- VIDAS® MEASLES IGG 60 TESTS (Ref. 30219) - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® MEASLES IGG 60 TESTS (Ref. 30219) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|--------------------------------|---------------------|---------------------|
| 30219 | 1008890110 | VIDAS® MEASLES IGG 60 TESTS | 16-Jun-22 | 14-Jun-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 4 related to FSCA 5333-1- VIDAS® RUB IGG II 60 TESTS (Ref. 30221) - -Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.

- Contact your local bioMérieux representative to order the replacement products when appropriate,



- As a reminder, please store the VIDAS® RUB IGG II 60 TESTS (Ref. 30221) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|----------------------------|---------------------|---------------------|
| 30221 | 1008566150 | VIDAS® RUB IGG II 60 TESTS | 2-Feb-22 | 12-Jan-22 |
| 30221 | 1008566660 | VIDAS® RUB IGG II 60 TESTS | 2-Feb-22 | 12-Jan-22 |
| 30221 | 1008611770 | VIDAS® RUB IGG II 60 TESTS | 1-Mar-22 | 9-Feb-22 |
| 30221 | 1008701600 | VIDAS® RUB IGG II 60 TESTS | 22-Apr-22 | 25-Mar-22 |
| 30221 | 1008809720 | VIDAS® RUB IGG II 60 TESTS | 15-Jun-22 | 17-May-22 |
| 30221 | 1008851780 | VIDAS® RUB IGG II 60 TESTS | 8-Jul-22 | 8-Jun-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 5 related to FSCA 5333-1- VIDAS® TOXO IGG AVIDITY 30 TESTS (Ref. 30222) - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.

- Contact your local bioMérieux representative to order the replacement products when appropriate,



- As a reminder, please store the VIDAS® TOXO IGG AVIDITY 30 TESTS (Ref. 30222) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|----------------------------------|---------------------|---------------------|
| 30222 | 1008721750 | VIDAS® TOXO IGG AVIDITY 30 TESTS | 23-Mar-22 | 10-Mar-22 |
| 30222 | 1008861600 | VIDAS® TOXO IGG AVIDITY 30 TESTS | 22-Jun-22 | 8-Jun-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 6 related to FSCA 5333-1- VIDAS® RUB IGG 60 TESTS (Ref. 30226) - Substrate error Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.

- Contact your local bioMérieux representative to order the replacement products when appropriate,



- As a reminder, please store the VIDAS® RUB IGG 60 TESTS (Ref. 30226) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|--------------------------------------|---------------------|---------------------|
| 30226 | 1008694980 | VIDAS® RUB IGG 60 TESTS (Ref. 30226) | 15-Apr-22 | 10-Mar-22 |
| 30226 | 1008700260 | VIDAS® RUB IGG 60 TESTS (Ref. 30226) | 15-Apr-22 | 10-Mar-22 |
| 30226 | 1008828980 | VIDAS® RUB IGG 60 TESTS (Ref. 30226) | 16-Jun-22 | 26-May-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 7 related to FSCA 5333-1- VIDAS® EBNA IGG 30 TESTS (Ref. 30235) - Substrate error Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.

- Contact your local bioMérieux representative to order the replacement products when appropriate,



- As a reminder, please store the VIDAS® EBNA IGG 30 TESTS (Ref. 30235) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|--------------------------|---------------------|---------------------|
| 30235 | 1008760410 | VIDAS® EBNA IGG 30 TESTS | 20-May-22 | 28-Apr-22 |
| 30235 | 1008908880 | VIDAS® EBNA IGG 30 TESTS | 2-Aug-22 | 13-Jul-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 8 related to FSCA 5333-1- VIDAS® EBV VCA.EA IGG 30 TESTS (Ref. 30236) - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.

- Contact your local bioMérieux representative to order the replacement products when appropriate,



- As a reminder, please store the VIDAS® EBV VCA.EA IGG 30 TESTS (Ref. 30236) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|--------------------------------|---------------------|---------------------|
| 30236 | 1008731540 | VIDAS® EBV VCA.EA IGG 30 TESTS | 4-May-22 | 7-Apr-22 |
| 30236 | 1008908700 | VIDAS® EBV VCA.EA IGG 30 TESTS | 2-Aug-22 | 13-Jul-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 9 related to FSCA 5333-1- VIDAS® EBV VCA IGM 30 TESTS (Ref. 30237) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.

- Contact your local bioMérieux representative to order the replacement products when appropriate,



- As a reminder, please store the VIDAS® EBV VCA IGM 30 TESTS (Ref. 30237) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible. We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|-----------------------------|---------------------|---------------------|
| 30237 | 1008733860 | VIDAS® EBV VCA IGM 30 TESTS | 15-Apr-22 | 25-Mar-22 |
| 30237 | 1008854070 | VIDAS® EBV VCA IGM 30 TESTS | 19-Jun-22 | 27-May-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 10 related to FSCA 5333-1- VIDAS® HBE/ANTI-HBE 30 TESTS (Ref. 30305) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.

- Contact your local bioMérieux representative to order the replacement products when appropriate,



- As a reminder, please store the VIDAS® HBE/ANTI-HBE 30 TESTS (Ref. 30305) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|---------------------------------|---------------------|---------------------|
| 30305 | 1008789580 | VIDAS® HBE.ANTI-HBE 30 TESTS | 19-May-22 | 19-Apr-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 11 related to FSCA 5333-1- VIDAS® ANTI-HAV TOTAL 30 TESTS (Ref. 30312) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.

- Contact your local bioMérieux representative to order the replacement products when appropriate,



- As a reminder, please store the VIDAS® ANTI-HAV TOTAL 30 TESTS (Ref. 30312) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|--------------------------------|---------------------|---------------------|
| 30312 | 1008589520 | VIDAS® ANTI-HAV TOTAL 30 TESTS | 16-Feb-22 | 25-Jan-22 |
| 30312 | 1008762100 | VIDAS® ANTI-HAV TOTAL 30 TESTS | 21-May-22 | 19-Apr-22 |
| 30312 | 1008920480 | VIDAS® ANTI-HAV TOTAL 30 TESTS | 17-Aug-22 | 24-Jul-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 12 related to FSCA 5333-1- VIDAS® LYME IGM 60 TESTS (Ref. 30319) - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1 and Table 2, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1 and 2, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1 and 2, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1 or Table 2.

- **For products with No remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify impacted lots in Table 1,
 - Stop using the listed impacted lots
 - Destroy the remaining lots in your inventory.



- **For products with remaining shelf life (taking into account the revised expiry date) in Table 2:**
 - Identify products listed in Table 2,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 2.
 - Continue product use until revised expiry date.
- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® LYME IGM 60 TESTS (Ref. 30319) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible. We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1: Product with NO remaining shelf life (taking into account the revised expiry date).

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|--------------|--------------------------|---------------------|---------------------|
| 30319 | 1008694220** | VIDAS® LYME IGM 60 TESTS | 13-Apr-22 | 2-Nov-21 |

** Lots under PSS (Product Stop Shipment)

Table 2 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|--------------------------|---------------------|---------------------|
| 30319 | 1008855250 | VIDAS® LYME IGM 60 TESTS | 6-Jul-22 | 18-Jan-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 13 related to FSCA 5333-1- VIDAS® LYME IGG 60 TESTS (Ref. 30320) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® LYME IGG 60 TESTS (Ref. 30320) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|--------------------------|---------------------|---------------------|
| 30320 | 1008800180 | VIDAS® LYME IGG 60 TESTS | 14-Jun-22 | 9-Jan-22 |
| 30320 | 1008900310 | VIDAS® LYME IGG 60 TESTS | 30-Jul-22 | 15-Feb-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 14 related to FSCA 5333-1- VIDAS® TSH 60 TESTS (Ref. 30400) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1 and Table 2, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1 and 2, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1 and 2, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1 or Table 2.

- **For products with No remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify impacted lots in Table 1,
 - Stop using the listed impacted lots
 - Destroy the remaining lots in your inventory.



- **For products with remaining shelf life (taking into account the revised expiry date) in Table 2:**
 - Identify products listed in Table 2,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 2.
 - Continue product use until revised expiry date.
- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® TSH 60 TESTS (Ref. 30400) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1: Product with NO remaining shelf life (taking into account the revised expiry date).

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|--------------|---------------------|---------------------|---------------------|
| 30400 | 1008581260** | VIDAS® TSH 60 TESTS | 17-Feb-22 | 6-Sep-21 |
| 30400 | 1008572270** | VIDAS® TSH 60 TESTS | 8-Feb-22 | 6-Sep-21 |

** Lots under PSS (Product Stop Shipment)

Table 2 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|---------------------|---------------------|---------------------|
| 30400 | 1008794370 | VIDAS® TSH 60 TESTS | 10-Jun-22 | 9-Jan-22 |
| 30400 | 1008784130 | VIDAS® TSH 60 TESTS | 29-May-22 | 9-Jan-22 |
| 30400 | 1008693180 | VIDAS® TSH 60 TESTS | 19-Apr-22 | 14-Nov-21 |
| 30400 | 1008708180 | VIDAS® TSH 60 TESTS | 21-Apr-22 | 27-Nov-21 |



| | | | | |
|-------|------------|---------------------|-----------|-----------|
| 30400 | 1008727020 | VIDAS® TSH 60 TESTS | 29-Apr-22 | 27-Nov-21 |
| 30400 | 1008800070 | VIDAS® TSH 60 TESTS | 14-Jun-22 | 9-Jan-22 |
| 30400 | 1008825270 | VIDAS® TSH 60 TESTS | 28-Jun-22 | 19-Jan-22 |
| 30400 | 1008840210 | VIDAS® TSH 60 TESTS | 2-Jul-22 | 24-Jan-22 |
| 30400 | 1008864740 | VIDAS® TSH 60 TESTS | 16-Jul-22 | 12-Feb-22 |
| 30400 | 1008886320 | VIDAS® TSH 60 TESTS | 26-Jul-22 | 22-Feb-22 |
| 30400 | 1008903990 | VIDAS® TSH 60 TESTS | 9-Aug-22 | 18-Mar-22 |
| 30400 | 1008923240 | VIDAS® TSH 60 TESTS | 17-Aug-22 | 7-Mar-22 |
| 30400 | 1008926570 | VIDAS® TSH 60 TESTS | 20-Aug-22 | 21-Mar-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 15 related to FSCA 5333-1- VIDAS® FT3 60 TESTS (Ref. 30402) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you**.

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® FT3 60 TESTS (Ref. 30402) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|---------------------|---------------------|---------------------|
| 30402 | 1008674380 | VIDAS® FT3 60 TESTS | 6-Apr-22 | 3-Mar-22 |
| 30402 | 1008584200 | VIDAS® FT3 60 TESTS | 16-Feb-22 | 26-Jan-22 |
| 30402 | 1008615890 | VIDAS® FT3 60 TESTS | 8-Mar-22 | 3-Feb-22 |
| 30402 | 1008704980 | VIDAS® FT3 60 TESTS | 19-Apr-22 | 25-Mar-22 |
| 30402 | 1008747690 | VIDAS® FT3 60 TESTS | 5-May-22 | 19-Apr-22 |
| 30402 | 1008775680 | VIDAS® FT3 60 TESTS | 28-May-22 | 4-May-22 |
| 30402 | 1008782770 | VIDAS® FT3 60 TESTS | 3-Jun-22 | 4-May-22 |
| 30402 | 1008845140 | VIDAS® FT3 60 TESTS | 2-Jul-22 | 8-Jun-22 |
| 30402 | 1008881770 | VIDAS® FT3 60 TESTS | 26-Jul-22 | 30-Jun-22 |
| 30402 | 1008914270 | VIDAS® FT3 60 TESTS | 12-Aug-22 | 30-Jun-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 16 related to FSCA 5333-1- VIDAS® T3 60 TESTS (Ref. 30403) - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you**.

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® T3 60 TESTS (Ref. 30403) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|--------------------|---------------------|---------------------|
| 30403 | 1008612240 | VIDAS® T3 60 TESTS | 2-Mar-22 | 25-Jan-22 |
| 30403 | 1008589770 | VIDAS® T3 60 TESTS | 18-Feb-22 | 25-Jan-22 |
| 30403 | 1008637510 | VIDAS® T3 60 TESTS | 17-Mar-22 | 23-Feb-22 |
| 30403 | 1008666330 | VIDAS® T3 60 TESTS | 25-Mar-22 | 3-Mar-22 |
| 30403 | 1008671650 | VIDAS® T3 60 TESTS | 2-Apr-22 | 3-Mar-22 |
| 30403 | 1008687730 | VIDAS® T3 60 TESTS | 9-Apr-22 | 22-Mar-22 |
| 30403 | 1008706040 | VIDAS® T3 60 TESTS | 22-Apr-22 | 25-Mar-22 |
| 30403 | 1008720580 | VIDAS® T3 60 TESTS | 4-May-22 | 7-Apr-22 |
| 30403 | 1008739140 | VIDAS® T3 60 TESTS | 10-May-22 | 19-Apr-22 |
| 30403 | 1008770970 | VIDAS® T3 60 TESTS | 3-Jun-22 | 28-Apr-22 |
| 30403 | 1008838990 | VIDAS® T3 60 TESTS | 2-Jul-22 | 8-Jun-22 |
| 30403 | 1008856820 | VIDAS® T3 60 TESTS | 9-Jul-22 | 14-Jun-22 |
| 30403 | 1008882160 | VIDAS® T3 60 TESTS | 21-Jul-22 | 20-Jun-22 |
| 30403 | 1008897120 | VIDAS® T3 60 TESTS | 5-Aug-22 | 30-Jun-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 17 related to FSCA 5333-1- VIDAS® T4 60 TESTS (Ref. 30404) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you**.

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Table 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® T4 60 TESTS (Ref. 30404) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|--------------------|---------------------|---------------------|
| 30404 | 1008679630 | VIDAS® T4 60 TESTS | 6-Apr-22 | 10-Mar-22 |
| 30404 | 1008598120 | VIDAS® T4 60 TESTS | 19-Feb-22 | 25-Jan-22 |
| 30404 | 1008735990 | VIDAS® T4 60 TESTS | 5-May-22 | 19-Apr-22 |
| 30404 | 1008815190 | VIDAS® T4 60 TESTS | 16-Jun-22 | 27-May-22 |
| 30404 | 1008843810 | VIDAS® T4 60 TESTS | 2-Jul-22 | 14-Jun-22 |
| 30404 | 1008905920 | VIDAS® T4 60 TESTS | 10-Aug-22 | 29-Jun-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 18 related to FSCA 5333-1- VIDAS® LH 60 TESTS (Ref. 30406) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you**.

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Table 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® LH 60 TESTS (Ref. 30406) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

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We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|--------------------|---------------------|---------------------|
| 30406 | 1008699160 | VIDAS® LH 60 TESTS | 19-Apr-22 | 17-Nov-21 |
| 30406 | 1008813780 | VIDAS® LH 60 TESTS | 14-Jun-22 | 9-Jan-22 |
| 30406 | 1008848620 | VIDAS® LH 60 TESTS | 1-Jul-22 | 5-Feb-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 19 related to FSCA 5333-1- VIDAS® FSH 60 TESTS (Ref. 30407) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Tables 1 and 2, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Tables 1 and 2, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1 and 2, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Tables 1 or 2.

- **For products with No remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify impacted lots in Table 1,
 - Stop using the listed impacted lots
 - Destroy the remaining lots in your inventory.



- **For products with remaining shelf life (taking into account the revised expiry date) in Table 2:**
 - Identify products listed in Table 2,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 2.
 - Continue product use until revised expiry date.
- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® FSH 60 TESTS (Ref. 30407) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1: Product with NO remaining shelf life (taking into account the revised expiry date).

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|--------------|---------------------|---------------------|---------------------|
| 30407 | 1008580750** | VIDAS® FSH 60 TESTS | 8-Feb-22 | 19-Sep-21 |
| 30407 | 1008660520** | VIDAS® FSH 60 TESTS | 22-Mar-22 | 18-Oct-21 |

** Lots under PSS (Product Stop Shipment)

Table 2 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|---------------------|---------------------|---------------------|
| 30407 | 1008730980 | VIDAS® FSH 60 TESTS | 4-May-22 | 30-Nov-21 |
| 30407 | 1008781450 | VIDAS® FSH 60 TESTS | 3-Jun-22 | 21-Dec-21 |
| 30407 | 1008794330 | VIDAS® FSH 60 TESTS | 3-Jun-22 | 21-Dec-21 |
| 30407 | 1008860760 | VIDAS® FSH 60 TESTS | 8-Jul-22 | 31-Jan-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 20 related to FSCA 5333-1- VIDAS® PROLACTINE 60 TESTS (Ref. 30410) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Tables 1 and 2, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Tables 1 and 2, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1 and 2, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Tables 1 or 2.

- **For products with No remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify impacted lots in Table 1,
 - Stop using the listed impacted lots
 - Destroy the remaining lots in your inventory.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 2:**
 - Identify products listed in Table 2,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 2.
 - Continue product use until revised expiry date.
- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® PROLACTINE 60 TESTS (Ref. 30410) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
 bioMérieux (Suisse) SA

Table 1: Product with NO remaining shelf life (taking into account the revised expiry date).

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|--------------|----------------------------|---------------------|---------------------|
| 30410 | 1008661910** | VIDAS® PROLACTINE 60 TESTS | 22-Mar-22 | 26-Oct-21 |

** Lots under PSS (Product Stop Shipment)

Table 2 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|----------------------------|---------------------|---------------------|
| 30410 | 1008753000 | VIDAS® PROLACTINE 60 TESTS | 18-May-22 | 19-Dec-21 |
| 30410 | 1008757170 | VIDAS® PROLACTINE 60 TESTS | 18-May-22 | 19-Dec-21 |
| 30410 | 1008829010 | VIDAS® PROLACTINE 60 TESTS | 28-Jun-22 | 25-Jan-22 |
| 30410 | 1008900440 | VIDAS® PROLACTINE 60 TESTS | 29-Jul-22 | 27-Feb-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 21 related to FSCA 5333-1- VIDAS® FERRITINE 60 TESTS (Ref. 30411) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you**.

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Table 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® FERRITINE 60 TESTS (Ref. 30411) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|---------------------------|---------------------|---------------------|
| 30411 | 1008747640 | VIDAS® FERRITINE 60 TESTS | 11-May-22 | 27-Nov-21 |
| 30411 | 1008748190 | VIDAS® FERRITINE 60 TESTS | 10-May-22 | 19-Dec-21 |
| 30411 | 1008755660 | VIDAS® FERRITINE 60 TESTS | 21-May-22 | 19-Dec-21 |
| 30411 | 1008757920 | VIDAS® FERRITINE 60 TESTS | 27-May-22 | 12-Dec-21 |
| 30411 | 1008793210 | VIDAS® FERRITINE 60 TESTS | 2-Jun-22 | 9-Jan-22 |
| 30411 | 1008809670 | VIDAS® FERRITINE 60 TESTS | 14-Jun-22 | 9-Jan-22 |
| 30411 | 1008812930 | VIDAS® FERRITINE 60 TESTS | 3-Jun-22 | 9-Jan-22 |
| 30411 | 1008815210 | VIDAS® FERRITINE 60 TESTS | 13-Jun-22 | 9-Jan-22 |
| 30411 | 1008816700 | VIDAS® FERRITINE 60 TESTS | 17-Jun-22 | 9-Jan-22 |
| 30411 | 1008821300 | VIDAS® FERRITINE 60 TESTS | 16-Jun-22 | 9-Jan-22 |
| 30411 | 1008828840 | VIDAS® FERRITINE 60 TESTS | 28-Jun-22 | 25-Jan-22 |
| 30411 | 1008838350 | VIDAS® FERRITINE 60 TESTS | 27-Jun-22 | 25-Jan-22 |
| 30411 | 1008845330 | VIDAS® FERRITINE 60 TESTS | 1-Jul-22 | 5-Feb-22 |
| 30411 | 1008854100 | VIDAS® FERRITINE 60 TESTS | 8-Jul-22 | 8-Feb-22 |



| | | | | |
|-------|------------|---------------------------|-----------|-----------|
| 30411 | 1008903380 | VIDAS® FERRITINE 60 TESTS | 2-Aug-22 | 22-Feb-22 |
| 30411 | 1008903160 | VIDAS® FERRITINE 60 TESTS | 1-Aug-22 | 7-Mar-22 |
| 30411 | 1008928580 | VIDAS® FERRITINE 60 TESTS | 20-Aug-22 | 21-Mar-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 22 related to FSCA 5333-1- VIDAS® AFP 60 TESTS (Ref. 30413) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Table 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® AFP 60 TESTS (Ref. 30413) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|---------------------|---------------------|---------------------|
| 30413 | 1008805600 | VIDAS® AFP 60 TESTS | 8-Jun-22 | 9-Jan-22 |
| 30413 | 1008921510 | VIDAS® AFP 60 TESTS | 11-Aug-22 | 7-Mar-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 23 related to FSCA 5333-1- VIDAS® IGE 60 TESTS (Ref. 30419) – - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Tables 1 and 2, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Tables 1 and 2, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1 and 2, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Tables 1 or 2.

- **For products with No remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify impacted lots in Table 1,
 - Stop using the listed impacted lots
 - Destroy the remaining lots in your inventory.



- **For products with remaining shelf life (taking into account the revised expiry date) in Table 2:**
 - Identify products listed in Table 2,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 2.
 - Continue product use until revised expiry date.
- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® IGE 60 TESTS (Ref. 30419) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1: Product with NO remaining shelf life (taking into account the revised expiry date).

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|--------------|---------------------|---------------------|---------------------|
| 30419 | 1008680690** | VIDAS® IGE 60 TESTS | 7-Apr-22 | 26-Oct-21 |

** Lots under PSS (Product Stop Shipment)

Table 2 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|---------------------|---------------------|---------------------|
| 30419 | 1008719960 | VIDAS® IGE 60 TESTS | 26-Apr-22 | 27-Nov-21 |
| 30419 | 1008777960 | VIDAS® IGE 60 TESTS | 27-May-22 | 27-Dec-21 |
| 30419 | 1008860770 | VIDAS® IGE 60 TESTS | 1-Jul-22 | 5-Feb-22 |
| 30419 | 1008882240 | VIDAS® IGE 60 TESTS | 15-Jul-22 | 5-Feb-22 |
| 30419 | 1008884980 | VIDAS® IGE 60 TESTS | 15-Jul-22 | 5-Feb-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 24 related to FSCA 5333-1- VIDAS® B2 MICROGLOBULIN 30 TESTS (Ref. 30420) - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Tables 1 and 2, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Tables 1 and 2, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1 and 2, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Tables 1 or 2.

- **For products with No remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify impacted lots in Table 1,
 - Stop using the listed impacted lots
 - Destroy the remaining lots in your inventory.



- **For products with remaining shelf life (taking into account the revised expiry date) in Table 2:**
 - Identify products listed in Table 2,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 2.
 - Continue product use until revised expiry date.
- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® B2 MICROGLOBULIN 30 TESTS (Ref. 30420) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1: Product with NO remaining shelf life (taking into account the revised expiry date).

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|--------------|--|---------------------|---------------------|
| 30420 | 1008595270** | VIDAS® B2 MICROGLOBULIN 30 TESTS | 12-Feb-22 | 20-Sep-21 |

** Lots under PSS (Product Stop Shipment)

Table 2 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|--|---------------------|---------------------|
| 30420 | 1008708170 | VIDAS® B2 MICROGLOBULIN 30 TESTS | 20-Apr-22 | 17-Nov-21 |
| 30420 | 1008856170 | VIDAS® B2 MICROGLOBULIN 30 TESTS | 7-Jul-22 | 6-Feb-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 25 related to FSCA 5333-1- VIDAS® CA 125 II 30 TESTS (Ref. 30426) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Tables 1 and 2, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Tables 1 and 2, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1 and 2, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Tables 1 or 2.

- **For products with No remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify impacted lots in Table 1,
 - Stop using the listed impacted lots
 - Destroy the remaining lots in your inventory.



- **For products with remaining shelf life (taking into account the revised expiry date) in Table 2:**
 - Identify products listed in Table 2,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 2.
 - Continue product use until revised expiry date.
- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® CA 125 II 30 TESTS (Ref. 30426) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1: Product with NO remaining shelf life (taking into account the revised expiry date).

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|--------------|---------------------------|---------------------|---------------------|
| 30426 | 1008594710** | VIDAS® CA 125 II 30 TESTS | 26-Jan-22 | 19-Sep-21 |
| 30426 | 1008684500** | VIDAS® CA 125 II 30 TESTS | 24-Mar-22 | 2-Nov-21 |

** Lots under PSS (Product Stop Shipment)

Table 2 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|---------------------------|---------------------|---------------------|
| 30426 | 1008908500 | VIDAS® CA 125 II 30 TESTS | 21-Jul-22 | 7-Mar-22 |
| 30426 | 1008884360 | VIDAS® CA 125 II 30 TESTS | 16-Jul-22 | 12-Feb-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 26 related to FSCA 5333-1- VIDAS® CA 19-9 30 TESTS (Ref. 30427) – - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Tables 1 and 2, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Tables 1 and 2, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1 and 2, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Tables 1 or 2.

- **For products with No remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify impacted lots in Table 1,
 - Stop using the listed impacted lots
 - Destroy the remaining lots in your inventory.



- **For products with remaining shelf life (taking into account the revised expiry date) in Table 2:**
 - Identify products listed in Table 2,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 2.
 - Continue product use until revised expiry date.
- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® CA 19-9 30 TESTS (Ref. 30427) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1: Product with NO remaining shelf life (taking into account the revised expiry date).

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|--------------|-------------------------|---------------------|---------------------|
| 30427 | 1008621790** | VIDAS® CA 19-9 30 TESTS | 17-Feb-22 | 4-Oct-21 |

** Lots under PSS (Product Stop Shipment)

Table 2 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|-------------------------|---------------------|---------------------|
| 30427 | 1008755840 | VIDAS® CA 19-9 30 TESTS | 11-May-22 | 12-Dec-21 |
| 30427 | 1008816890 | VIDAS® CA 19-9 30 TESTS | 25-May-22 | 19-Jan-22 |
| 30427 | 1008928460 | VIDAS® CA 19-9 30 TESTS | 8-Jul-22 | 6-Feb-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 27 related to FSCA 5333-1- VIDAS® TPSA 60 TESTS (Ref. 30428) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Tables 1 and 2, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Tables 1 and 2, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1 and 2, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Tables 1 or 2.

- **For products with No remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify impacted lots in Table 1,
 - Stop using the listed impacted lots
 - Destroy the remaining lots in your inventory.



- **For products with remaining shelf life (taking into account the revised expiry date) in Table 2:**
 - Identify products listed in Table 2,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 2.
 - Continue product use until revised expiry date.
- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® TPSA 60 TESTS (Ref. 30428) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible. We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1: Product with NO remaining shelf life (taking into account the revised expiry date).

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|--------------|----------------------|---------------------|---------------------|
| 30428 | 1008678810** | VIDAS® TPSA 60 TESTS | 26-Mar-22 | 2-Nov-21 |
| 30428 | 1008616040** | VIDAS® TPSA 60 TESTS | 22-Feb-22 | 28-Sep-21 |

** Lots under PSS (Product Stop Shipment)

Table 2 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|----------------------|---------------------|---------------------|
| 30428 | 1008757280 | VIDAS® TPSA 60 TESTS | 4-May-22 | 12-Dec-21 |
| 30428 | 1008705160 | VIDAS® TPSA 60 TESTS | 14-Apr-22 | 17-Nov-21 |
| 30428 | 1008830760 | VIDAS® TPSA 60 TESTS | 22-Jun-22 | 24-Jan-22 |
| 30428 | 1008842800 | VIDAS® TPSA 60 TESTS | 23-Jun-22 | 31-Jan-22 |
| 30428 | 1008930910 | VIDAS® TPSA 60 TESTS | 6-Aug-22 | 18-Mar-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 28 related to FSCA 5333-1- VIDAS® CA 15-3 30 TESTS (Ref. 30429) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Tables 1 and 2, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Tables 1 and 2, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1 and 2, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1 or 2.

- **For products with No remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify impacted lots in Table 1,
 - Stop using the listed impacted lots
 - Destroy the remaining lots in your inventory.



- **For products with remaining shelf life (taking into account the revised expiry date) in Table 2:**
 - Identify products listed in Table 2,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 2.
 - Continue product use until revised expiry date.
- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® CA 15-3 30 TESTS (Ref. 30429) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1: Product with NO remaining shelf life (taking into account the revised expiry date).

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|--------------|-------------------------|---------------------|---------------------|
| 30429 | 1008686710** | VIDAS® CA 15-3 30 TESTS | 9-Apr-22 | 2-Nov-21 |

** Lots under PSS (Product Stop Shipment)

Table 2 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|-------------------------|---------------------|---------------------|
| 30429 | 1008835210 | VIDAS® CA 15-3 30 TESTS | 29-Jun-22 | 18-Jan-22 |
| 30429 | 1008868770 | VIDAS® CA 15-3 30 TESTS | 20-Jul-22 | 6-Feb-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 29 related to FSCA 5333-1- VIDAS® ESTRADIOL II 60 TESTS (Ref. 30431) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® ESTRADIOL II 60 TESTS (Ref. 30431) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|------------------------------|---------------------|---------------------|
| 30431 | 1008671380 | VIDAS® ESTRADIOL II 60 TESTS | 29-Mar-22 | 3-Mar-22 |
| 30431 | 1008757870 | VIDAS® ESTRADIOL II 60 TESTS | 18-May-22 | 19-Apr-22 |
| 30431 | 1008856950 | VIDAS® ESTRADIOL II 60 TESTS | 22-Jun-22 | 27-May-22 |
| 30431 | 1008872270 | VIDAS® ESTRADIOL II 60 TESTS | 19-Jul-22 | 14-Jun-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 30 related to FSCA 5333-1- VIDAS® VWF 30 TESTS (Ref. 30436) – - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® VWF 30 TESTS (Ref. 30436) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|---------------------|---------------------|---------------------|
| 30436 | 1008622630 | VIDAS® VWF 30 TESTS | 9-Mar-22 | 9-Feb-22 |
| 30436 | 1008939670 | VIDAS® VWF 30 TESTS | 10-Aug-22 | 29-Jun-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 31 related to FSCA 5333-1- VIDAS® FPSA 30 TESTS (Ref. 30440) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Tables 1 and 2, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Tables 1 and 2, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1 and 2, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Tables 1 or 2.

- **For products with No remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify impacted lots in Table 1,
 - Stop using the listed impacted lots
 - Destroy the remaining lots in your inventory.



- **For products with remaining shelf life (taking into account the revised expiry date) in Table 2:**
 - Identify products listed in Table 2,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 2.
 - Continue product use until revised expiry date.
- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® FPSA 30 TESTS (Ref. 30440) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1: Product with NO remaining shelf life (taking into account the revised expiry date).

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|--------------|----------------------|---------------------|---------------------|
| 30440 | 1008654860** | VIDAS® FPSA 30 TESTS | 18-Mar-22 | 26-Oct-21 |
| 30440 | 1008699150** | VIDAS® FPSA 30 TESTS | 14-Apr-22 | 2-Nov-21 |
| 30440 | 1008605070** | VIDAS® FPSA 30 TESTS | 16-Feb-22 | 28-Sep-21 |

** Lots under PSS (Product Stop Shipment)

Table 2 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|----------------------|---------------------|---------------------|
| 30440 | 1008763720 | VIDAS® FPSA 30 TESTS | 3-May-22 | 12-Dec-21 |
| 30440 | 1008852160 | VIDAS® FPSA 30 TESTS | 1-Jul-22 | 31-Jan-22 |
| 30440 | 1008931040 | VIDAS® FPSA 30 TESTS | 9-Aug-22 | 18-Mar-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 32 related to FSCA 5333-1- VIDAS® TSH 3 60 TESTS (Ref. 30441) – - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Table 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® TSH 3 60 TESTS (Ref. 30441) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|-----------------------|---------------------|---------------------|
| 30441 | 1008843310 | VIDAS® TSH 3 60 TESTS | 25-Jun-22 | 5-Feb-22 |
| 30441 | 1008843340 | VIDAS® TSH 3 60 TESTS | 25-Jun-22 | 5-Feb-22 |
| 30441 | 1008891670 | VIDAS® TSH 3 60 TESTS | 21-Jul-22 | 27-Feb-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 33 related to FSCA 5333-1- VIDAS® CEA (S) 60 TESTS (Ref. 30453) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1 and Table 2, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1 and 2, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1 and 2, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1 or Table 2.

- **For products with No remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify impacted lots in Table 1,
 - Stop using the listed impacted lots
 - Destroy the remaining lots in your inventory.



- **For products with remaining shelf life (taking into account the revised expiry date) in Table 2:**
 - Identify products listed in Table 2,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 2.
 - Continue product use until revised expiry date.
- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® CEA (S) 60 TESTS (Ref. 30453) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1: Product with NO remaining shelf life (taking into account the revised expiry date).

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|--------------|-------------------------|---------------------|---------------------|
| 30453 | 1008626550** | VIDAS® CEA (S) 60 TESTS | 8-Mar-22 | 4-Oct-21 |

** Lots under PSS (Product Stop Shipment)

Table 2 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|-------------------------|---------------------|---------------------|
| 30453 | 1008705700 | VIDAS® CEA (S) 60 TESTS | 20-Apr-22 | 17-Nov-21 |
| 30453 | 1008827310 | VIDAS® CEA (S) 60 TESTS | 21-Jun-22 | 19-Jan-22 |
| 30453 | 1008888060 | VIDAS® CEA (S) 60 TESTS | 26-Jul-22 | 22-Feb-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 34 related to FSCA 5333-1- VIDAS® FT4N 60 TESTS (Ref. 30459) – - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® FT4N 60 TESTS (Ref. 30459) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|----------------------|---------------------|---------------------|
| 30459 | 1008576580 | VIDAS® FT4N 60 TESTS | 10-Feb-22 | 25-Jan-22 |
| 30459 | 1008622440 | VIDAS® FT4N 60 TESTS | 5-Mar-22 | 9-Feb-22 |
| 30459 | 1008652140 | VIDAS® FT4N 60 TESTS | 22-Mar-22 | 23-Feb-22 |
| 30459 | 1008589800 | VIDAS® FT4N 60 TESTS | 10-Feb-22 | 25-Jan-22 |
| 30459 | 1008695620 | VIDAS® FT4N 60 TESTS | 13-Apr-22 | 25-Mar-22 |
| 30459 | 1008714720 | VIDAS® FT4N 60 TESTS | 23-Apr-22 | 4-Apr-22 |
| 30459 | 1008791570 | VIDAS® FT4N 60 TESTS | 7-Jun-22 | 17-May-22 |
| 30459 | 1008817980 | VIDAS® FT4N 60 TESTS | 14-Jun-22 | 17-May-22 |
| 30459 | 1008827280 | VIDAS® FT4N 60 TESTS | 22-Jun-22 | 1-Jun-22 |
| 30459 | 1008852180 | VIDAS® FT4N 60 TESTS | 8-Jul-22 | 14-Jun-22 |
| 30459 | 1008888200 | VIDAS® FT4N 60 TESTS | 27-Jul-22 | 30-Jun-22 |
| 30459 | 1008918000 | VIDAS® FT4N 60 TESTS | 12-Aug-22 | 27-Jul-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 35 related to FSCA 5333-1- VIDAS® ANTI-TPO 30 T (Ref. 30461) – - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® ANTI-TPO 30 T (Ref. 30461) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|----------------------|---------------------|---------------------|
| 30461 | 1008772240 | VIDAS® ANTI-TPO 30 T | 21-May-23 | 26-Apr-22 |
| 30461 | 1008845910 | VIDAS® ANTI-TPO 30 T | 24-Jun-23 | 2-Jun-22 |
| 30461 | 1008857460 | VIDAS® ANTI-TPO 30 T | 5-Jul-23 | 13-Jun-22 |
| 30461 | 1008926950 | VIDAS® ANTI-TPO 30 T | 5-Aug-23 | 28-Jul-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 36 related to FSCA 5333-1- VIDAS® ANTI-TG 30 T (Ref. 30462) – - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® ANTI-TG 30 T (Ref. 30462) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|---------------------|---------------------|---------------------|
| 30462 | 1008748270 | VIDAS® ANTI-TG 30 T | 2-Aug-22 | 4-Apr-22 |
| 30462 | 1008770010 | VIDAS® ANTI-TG 30 T | 15-Aug-22 | 9-May-22 |
| 30462 | 1008777840 | VIDAS® ANTI-TG 30 T | 31-Aug-22 | 26-Apr-22 |
| 30462 | 1008862600 | VIDAS® ANTI-TG 30 T | 22-Sep-22 | 13-Jun-22 |
| 30462 | 1008866080 | VIDAS® ANTI-TG 30 T | 3-Oct-22 | 23-Jun-22 |
| 30462 | 1008904010 | VIDAS® ANTI-TG 30 T | 20-Oct-22 | 5-Jul-22 |
| 30462 | 1008924530 | VIDAS® ANTI-TG 30 T | 3-Nov-22 | 5-Jul-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 37 related to FSCA 5333-1- VIDAS® 25-OH VITAMINE D TOTAL 60T (Ref. 30463) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.

- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® 25-OH VITAMINE D TOTAL 60T (Ref. 30463) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
 bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|-------|------------|-----------------------------------|---------------------|---------------------|
| 30463 | 1008764090 | VIDAS® 25-OH VITAMINE D TOTAL 60T | 25-Aug-22 | 19-Apr-22 |
| 30463 | 1008609060 | VIDAS® 25-OH VITAMINE D TOTAL 60T | 25-May-22 | 9-Feb-22 |
| 30463 | 1008676430 | VIDAS® 25-OH VITAMINE D TOTAL 60T | 30-Jun-22 | 3-Mar-22 |
| 30463 | 1008693940 | VIDAS® 25-OH VITAMINE D TOTAL 60T | 18-Jul-22 | 10-Mar-22 |
| 30463 | 1008717350 | VIDAS® 25-OH VITAMINE D TOTAL 60T | 28-Jul-22 | 7-Apr-22 |
| 30463 | 1008747560 | VIDAS® 25-OH VITAMINE D TOTAL 60T | 9-Aug-22 | 19-Apr-22 |
| 30463 | 1008772230 | VIDAS® 25-OH VITAMINE D TOTAL 60T | 31-Aug-22 | 28-Apr-22 |
| 30463 | 1008776690 | VIDAS® 25-OH VITAMINE D TOTAL 60T | 25-Aug-22 | 19-Apr-22 |
| 30463 | 1008796740 | VIDAS® 25-OH VITAMINE D TOTAL 60T | 5-Sep-22 | 17-May-22 |

| | | | | |
|-------|------------|---|-----------|-----------|
| 30463 | 1008841360 | VIDAS® 25-OH VITAMINE D TOTAL 60T | 29-Sep-22 | 8-Jun-22 |
| 30463 | 1008869580 | VIDAS® 25-OH VITAMINE D TOTAL 60T | 18-Oct-22 | 20-Jun-22 |
| 30463 | 1008892600 | VIDAS® 25-OH VITAMINE D TOTAL 60T | 25-Oct-22 | 30-Jun-22 |
| 30463 | 1008914140 | VIDAS® 25-OH VITAMINE D TOTAL 60T | 15-Nov-22 | 24-Jul-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 38 related to FSCA 5333-1- VIDAS® TESTOSTERONE II 30 TESTS (Ref. 414320) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® TESTOSTERONE II 30 TESTS (Ref. 414320) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|--------|------------|---------------------------------------|---------------------|---------------------|
| 414320 | 1008755640 | VIDAS® TESTOSTERONE II 30 TESTS | 15-Aug-22 | 26-Apr-22 |
| 414320 | 1008815760 | VIDAS® TESTOSTERONE II 30 TESTS | 9-Sep-22 | 2-Jun-22 |
| 414320 | 1008862630 | VIDAS® TESTOSTERONE II 30 TESTS | 3-Oct-22 | 13-Jun-22 |
| 414320 | 1008892260 | VIDAS® TESTOSTERONE II 30 TESTS | 21-Oct-22 | 23-Jun-22 |
| 414320 | 1008905890 | VIDAS® TESTOSTERONE II 30 TESTS | 26-Oct-22 | 5-Jul-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 39 related to FSCA 5333-1- VIDAS® LYME IGM II 60 TESTS (Ref. 416436) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1 and Table 2, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1 and 2, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1 and 2, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1 or Table 2.

- **For products with No remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify impacted lots in Table 1,
 - Stop using the listed impacted lots
 - Destroy the remaining lots in your inventory.



- **For products with remaining shelf life (taking into account the revised expiry date) in Table 2:**
 - Identify products listed in Table 2,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 2.
 - Continue product use until revised expiry date.
- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® LYME IGM II 60 TESTS (Ref. 416436) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1: Product with NO remaining shelf life (taking into account the revised expiry date).

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|--------|--------------|-----------------------------|---------------------|---------------------|
| 416436 | 1008636750** | VIDAS® LYME IGM II 60 TESTS | 9-Jun-22 | 20-Sep-21 |

** Lots under PSS (Product Stop Shipment)

Table 2 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|--------|------------|-----------------------------|---------------------|---------------------|
| 416436 | 1008768920 | VIDAS® LYME IGM II 60 TESTS | 15-Aug-22 | 12-Dec-21 |
| 416436 | 1008857410 | VIDAS® LYME IGM II 60 TESTS | 3-Oct-22 | 24-Jan-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 40 related to FSCA 5333-1- VIDAS® AMH 30 TESTS (Ref. 417011) – - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you**.

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® AMH 30 TESTS (Ref. 417011) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|--------|------------|---------------------|---------------------|---------------------|
| 417011 | 1008605020 | VIDAS® AMH 30 TESTS | 28-Aug-22 | 9-Feb-22 |
| 417011 | 1008667580 | VIDAS® AMH 30 TESTS | 26-Sep-22 | 3-Mar-22 |
| 417011 | 1008730610 | VIDAS® AMH 30 TESTS | 31-Oct-22 | 7-Apr-22 |
| 417011 | 1008854280 | VIDAS® AMH 30 TESTS | 3-Jan-23 | 13-Jun-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 41 related to FSCA 5333-1- VIDAS® LYME IGG II 60 TESTS (Ref. 417401) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® LYME IGG II 60 TESTS (Ref. 417401) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|--------|------------|-----------------------------|---------------------|---------------------|
| 417401 | 1008714880 | VIDAS® LYME IGG II 60 TESTS | 25-Jul-22 | 30-Nov-21 |
| 417401 | 1008824220 | VIDAS® LYME IGG II 60 TESTS | 12-Sep-22 | 9-Jan-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 42 related to FSCA 5333-1- VIDAS® HEV IGG 30T (Ref. 418116) – - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you**.

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® HEV IGG 30T (Ref. 418116) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|--------|------------|--------------------|---------------------|---------------------|
| 418116 | 1008684310 | VIDAS® HEV IGG 30T | 3-Oct-22 | 10-Mar-22 |
| 418116 | 1008831640 | VIDAS® HEV IGG 30T | 25-Dec-22 | 26-May-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 43 related to FSCA 5333-1- VIDAS® VIDAS PTH (1-84) 30T (Ref. 422010) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you**.

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® VIDAS PTH (1-84) 30T (Ref. 422010) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|--------|------------|-----------------------------|---------------------|---------------------|
| 422010 | 1008572010 | VIDAS® VIDAS PTH (1-84) 30T | 8-Feb-22 | 12-Jan-22 |
| 422010 | 1008690000 | VIDAS® VIDAS PTH (1-84) 30T | 1-Apr-22 | 10-Mar-22 |
| 422010 | 1008842830 | VIDAS® VIDAS PTH (1-84) 30T | 24-Jun-22 | 26-May-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 44 related to FSCA 5333-1- VIDAS® ANTI-DENGUE IGG 60 TESTS (Ref. 423079) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you**.

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® ANTI-DENGUE IGG 60 TESTS (Ref. 423079) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|--------|------------|---------------------------------|---------------------|---------------------|
| 423079 | 1008693240 | VIDAS® ANTI-DENGUE IGG 60 TESTS | 16-Oct-22 | 10-Mar-22 |
| 423079 | 1008832050 | VIDAS® ANTI-DENGUE IGG 60 TESTS | 25-Dec-22 | 26-May-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 45 related to FSCA 5333-1- VIDAS® TB-IGRA 60 TESTS (Ref. 423111) – - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® TB-IGRA 60 TESTS (Ref. 423111) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|--------|------------|-------------------------|---------------------|---------------------|
| 423111 | 1008779880 | VIDAS® TB-IGRA 60 TESTS | 28-Nov-22 | 28-Apr-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 46 related to FSCA 5333-1- VIDAS® SARS-COV-2 IgM (9COM) 60T (Ref. 423833) - - Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you**.

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS[®] SARS-COV-2 IgM (9COM) 60T (Ref. 423833) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|--------|------------|--|---------------------|---------------------|
| 423833 | 1008710940 | VIDAS [®] SARS-COV-2 IgM (9COM) 60T | 4-Mar-22 | 30-Nov-21 |
| 423833 | 1008771420 | VIDAS [®] SARS-COV-2 IgM (9COM) 60T | 18-May-22 | 31-Dec-21 |
| 423833 | 1008781210 | VIDAS [®] SARS-COV-2 IgM (9COM) 60T | 31-May-22 | 31-Dec-21 |
| 423833 | 1008787590 | VIDAS [®] SARS-COV-2 IgM (9COM) 60T | 7-Jun-22 | 31-Dec-21 |
| 423833 | 1008812250 | VIDAS [®] SARS-COV-2 IgM (9COM) 60T | 16-Jun-22 | 18-Jan-22 |
| 423833 | 1008843370 | VIDAS [®] SARS-COV-2 IgM (9COM) 60T | 28-Jun-22 | 31-Jan-22 |
| 423833 | 1008851220 | VIDAS [®] SARS-COV-2 IgM (9COM) 60T | 5-Jul-22 | 24-Jan-22 |
| 423833 | 1008886980 | VIDAS [®] SARS-COV-2 IgM (9COM) 60T | 22-Jul-22 | 22-Feb-22 |
| 423833 | 1008918390 | VIDAS [®] SARS-COV-2 IgM (9COM) 60T | 16-Aug-22 | 18-Mar-22 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 47 related to FSCA 5333-1- VIDAS® SARS-COV-2 IgG (9COG) 60T (Ref. 423834) - -Substrate error - Potential delayed results without medical impact

Dear bioMérieux Customer,

This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1 and Table 2, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications.

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1 and 2, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results. For impacted VIDAS® immuno-assay products listed in Tables 1 and 2, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1 or Table 2.

- **For products with No remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify impacted lots in Table 1,
 - Stop using the listed impacted lots
 - Destroy the remaining lots in your inventory.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 2:**
 - Identify products listed in Table 2,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 2.
 - Continue product use until revised expiry date.
- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® SARS-COV-2 IgG (9COG) 60T (Ref. 423834) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible. We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
 bioMérieux (Suisse) SA

Table 1: Product with NO remaining shelf life (taking into account the revised expiry date).

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|--------|--------------|----------------------------------|---------------------|---------------------|
| 423834 | 1008630790** | VIDAS® SARS-COV-2 IgG (9COG) 60T | 23-Feb-22 | 4-Oct-21 |
| 423834 | 1008671420** | VIDAS® SARS-COV-2 IgG (9COG) 60T | 29-Mar-22 | 26-Oct-21 |
| 423834 | 1008674550** | VIDAS® SARS-COV-2 IgG (9COG) 60T | 6-Apr-22 | 26-Oct-21 |
| 423834 | 1008685700** | VIDAS® SARS-COV-2 IgG (9COG) 60T | 14-Apr-22 | 2-Nov-21 |

** Lots under PSS (Product Stop Shipment)

Table 2 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|--------|------------|----------------------------------|---------------------|---------------------|
| 423834 | 1008714650 | VIDAS® SARS-COV-2 IgG (9COG) 60T | 28-Apr-22 | 17-Nov-21 |
| 423834 | 1008730600 | VIDAS® SARS-COV-2 IgG (9COG) 60T | 30-Apr-22 | 30-Nov-21 |
| 423834 | 1008747950 | VIDAS® SARS-COV-2 IgG (9COG) 60T | 11-May-22 | 12-Dec-21 |
| 423834 | 1008750690 | VIDAS® SARS-COV-2 IgG (9COG) 60T | 17-May-22 | 12-Dec-21 |



IMPORTANT, URGENT FIELD SAFETY NOTICE

Attachment 48 related to FSCA 5333-1- VIDAS® SARS-COV-2 IgG II (Ref. 424114) – - Substrate error - Potential delayed results without medical impact

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This FSCA #5333-1 is a follow-up of FSCA #5333 (released on 22-Sep-2021). FSCA #5333-1 concerns all lots of clinical VIDAS® immuno-assay product references for which a delayed result **has no medical impact but can lead to a major inconvenience for you.**

Our records indicate that your laboratory received one/several of the lots indicated in Table 1, below.

Description of the issue:

Based on an unusual rate of complaints from the field for “substrate error” leading to a potential delayed results on different VIDAS® references tested on all VIDAS® systems, bioMérieux initiated an investigation.

The substrate error issue was confirmed on all lots of VIDAS® Immuno-Assays manufactured with substrate batches using the common lot of raw material (4-MUP). The only potential risk associated with the substrate error is a delayed result as you need to run another test. There is no risk of false results. The investigation demonstrates that the issue occurs over the product shelf-life. The analyzes of the kinetic model allowed to define a revised expiry date for each impacted lot of VIDAS® Immuno-Assays finished products. When used until the revised expiry date, the product continues to perform per its registered performance specifications

Therefore, corrective actions involving a revised expiration date for all lots of clinical VIDAS® Immuno-assay products are required to ensure that the specified products will continue to perform per their labelled performance specifications.

Indeed, even if there is no medical impact of a delayed result when using the impacted lots of clinical VIDAS® Immuno-assay products listed in Table 1, as a conservative approach and to avoid a major inconvenience at your level, bioMérieux has decided to implement a Field Safety Corrective Action (FSCA #5333-1).

Impact to Customer/Patient:

In case of substrate error, there is a potential for delayed results. There is no risk of false results.

For impacted VIDAS® immuno-assay products listed in Tables 1, there is no medical impact of a delayed result but the issue can lead to a major inconvenience.

Required actions:

We request you to take the following actions at this time:

Following receipt of this Urgent Field Safety Notice, check inventory to determine if lots are listed in Table 1.

- **For products with remaining shelf life (taking into account the revised expiry date) in Table 1:**
 - Identify products listed in Table 1,
 - Update product expiry date per your internal procedures of the remaining usable lots to meet the new revised expiry date specified in Table 1.
 - Continue product use until revised expiry date.



- Contact your local bioMérieux representative to order the replacement products when appropriate,
- As a reminder, please store the VIDAS® SARS-COV-2 IgG II (Ref. 424114) at 2-8°C as described in Product Instructions for Use.
- Please distribute this information to all appropriate personnel in your laboratory, retain a copy in your files, and forward this information to all parties that may use this product, including others to whom you may have transferred our product.
- Complete the Acknowledgement Form in Attachment A and return it to your local bioMérieux representative by email (ch_support@biomerieux.com) to confirm receipt of this notice.

bioMérieux is committed to providing our customers with the highest quality product possible.

We sincerely apologize for any inconvenience that this may have caused you. If you require additional assistance or have any questions, please contact your Scientific Customer Service of bioMérieux at 022 906 57 96.

Sincerely
bioMérieux (Suisse) SA

Table 1 : Product with remaining shelf life (taking into account the revised expiry date)

| REF | Lot # | Product Name | Current Expiry date | Revised Expiry date |
|--------|------------|--------------------------|---------------------|---------------------|
| 424114 | 1008693260 | VIDAS® SARS-COV-2 IgG II | 19-Apr-22 | 25-Mar-22 |
| 424114 | 1008737650 | VIDAS® SARS-COV-2 IgG II | 3-May-22 | 19-Apr-22 |
| 424114 | 1008766600 | VIDAS® SARS-COV-2 IgG II | 25-May-22 | 19-Apr-22 |
| 424114 | 1008768870 | VIDAS® SARS-COV-2 IgG II | 31-May-22 | 19-Apr-22 |
| 424114 | 1008793180 | VIDAS® SARS-COV-2 IgG II | 8-Jun-22 | 8-May-22 |
| 424114 | 1008799340 | VIDAS® SARS-COV-2 IgG II | 7-Jun-22 | 26-May-22 |
| 424114 | 1008806170 | VIDAS® SARS-COV-2 IgG II | 11-Jun-22 | 26-May-22 |
| 424114 | 1008812180 | VIDAS® SARS-COV-2 IgG II | 10-Jun-22 | 26-May-22 |
| 424114 | 1008821130 | VIDAS® SARS-COV-2 IgG II | 21-Jun-22 | 26-May-22 |
| 424114 | 1008826440 | VIDAS® SARS-COV-2 IgG II | 20-Jun-22 | 26-May-22 |
| 424114 | 1008840470 | VIDAS® SARS-COV-2 IgG II | 1-Jul-22 | 8-Jun-22 |
| 424114 | 1008845260 | VIDAS® SARS-COV-2 IgG II | 5-Jul-22 | 1-Jun-22 |
| 424114 | 1008850980 | VIDAS® SARS-COV-2 IgG II | 6-Jul-22 | 1-Jun-22 |
| 424114 | 1008859210 | VIDAS® SARS-COV-2 IgG II | 12-Jul-22 | 14-Jun-22 |



| | | | | |
|--------|------------|-----------------------------|-----------|-----------|
| 424114 | 1008881730 | VIDAS® SARS-COV-2 IgG II | 19-Jul-22 | 23-Jun-22 |
| 424114 | 1008915730 | VIDAS® SARS-COV-2 IgG II | 9-Aug-22 | 5-Jul-22 |