



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Prepared by QA Committee		
Issued by: Laboratory Manager	<b>Revision Date:6/12/2025</b>	
Approved by Laboratory Director: Microbiologist-in-Chief	<b>Next Review Date:6/12/2027</b>	

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
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## Procedure of MALDI


### *VITEK MS Slide spot preparation*

### **Safety Precautions**

Wear protective gloves/protective clothing/eye protection/face protection

Before beginning to apply spots on a Vitek slide ensure slide is clean and reagents (matrix and formic acid) are at room temperature.

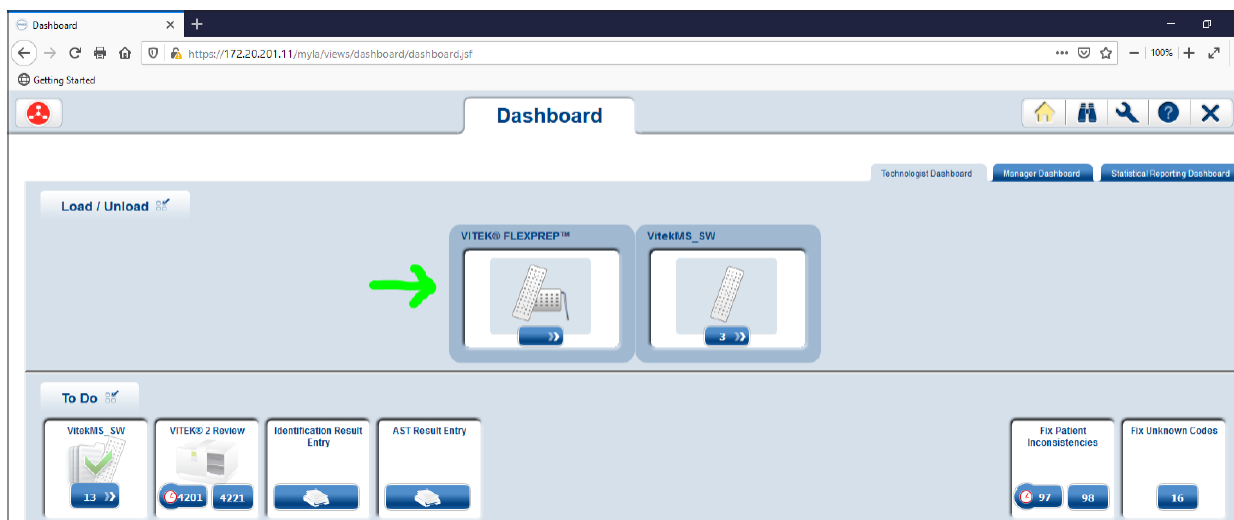
1. Label the [Vitek MS Worksheet](#) or [Vitek MS Worksheet with Controls](#) with the date and slide number
2. Apply a fresh isolate (24 hrs) of E. coli ATCC 8739 control in the central circle of the slide using a [sterile toothpick](#) for each acquisition group used.
3. Add 1ul of Matrix. Do not invert the matrix vial. Do not touch the crystals on the bottom of the vial with pipette tip
4. Ensure plates to be tested match the accompanying LIS label
5. Place one specimen barcode label on the work sheet
6. Apply a thin even layer of organism in each circle of the slide by using a [sterile toothpick](#)
7. Add 1ul of Matrix. Do not invert the matrix vial. Do not touch the crystals on the bottom of the vial with pipette tip
8. For Yeast, add 0.5ul of formic acid first, wait until it crystallizes, then add 1ul matrix,

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
### ***VITEK MS Prep Station***

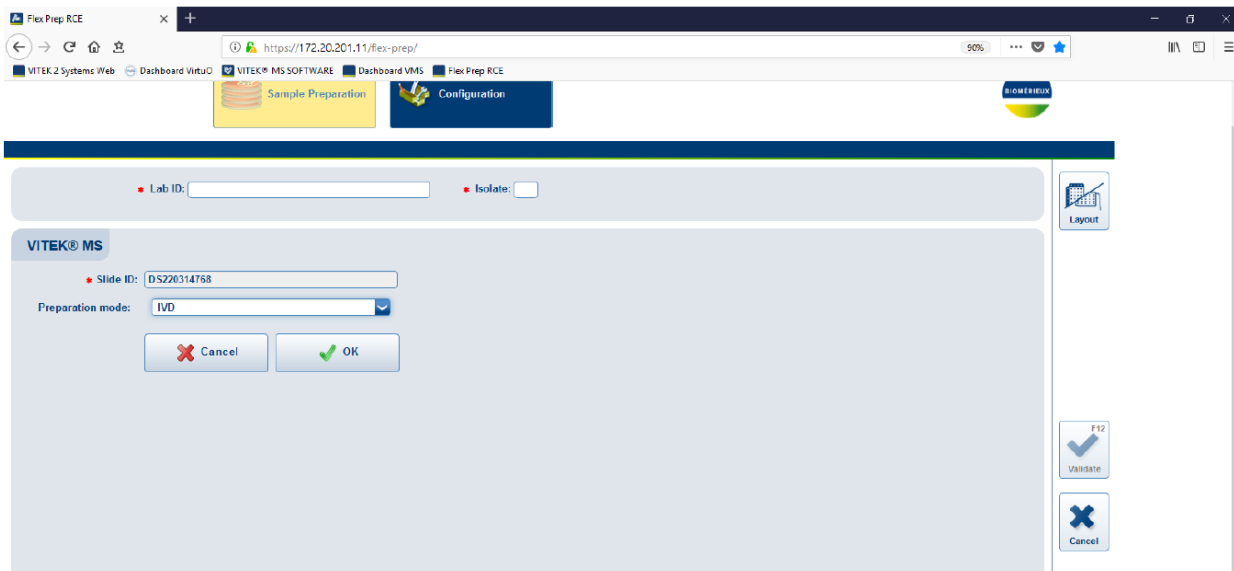
1. Double click the MYLA icon or also accessible via the Internet Address (192.168.168.241) in an Internet Browser window.
2. Enter User name
3. Enter Password

**Note:** Check Application Login Information in Microbiology Internal Manual or bench for posting of user name and current password

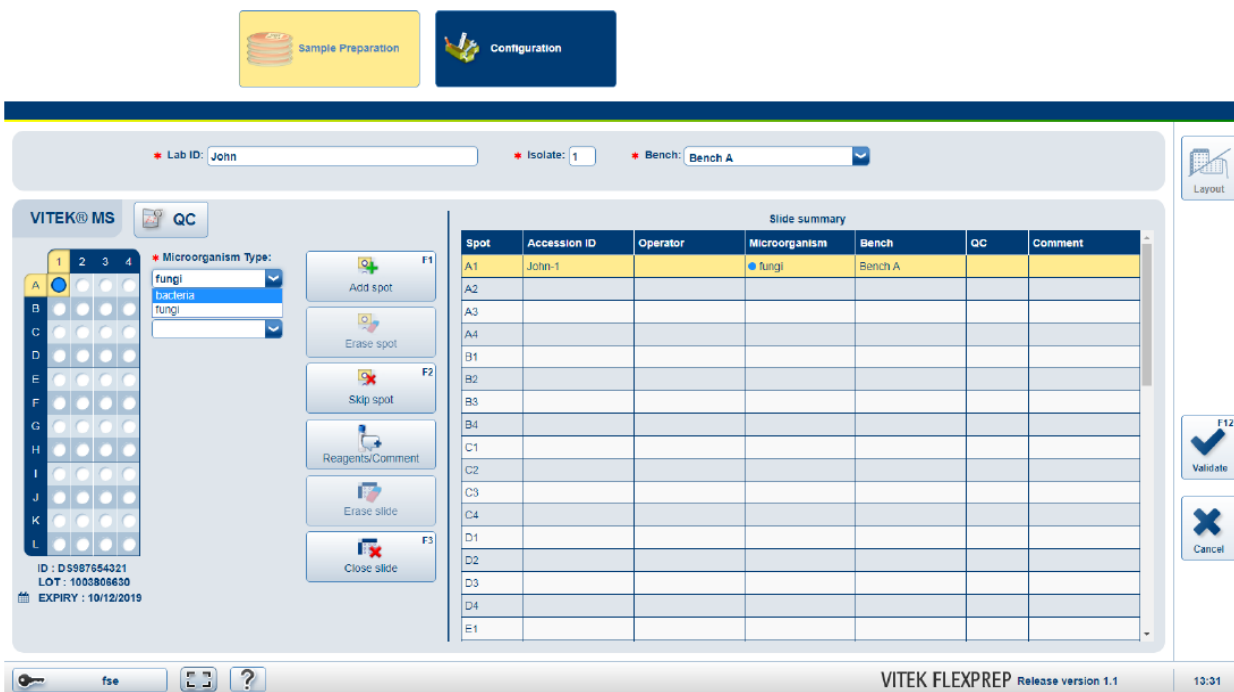


4. Click Vitek Flexprep icon (if needed click Layout button on right to toggle between Vitek XL and Vitek MS.)
5. Scan your Slide into the Slide ID field and select OK.

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6. Select your Bench ID from the dropdown next to "isolate".




7. Scan the specimen barcode on your worksheet in the Lab id# field

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- Ensure isolate number is correct – automatically increases each time barcode is entered on slide
- Bacteria is the default option
- For yeast spots, choose Fungi option

*Note: You may optionally duplicate the information for the next spot on the slide if you have made 2 spots for the same organism (ie. The same organism you are trying to ID with the same specimen **and** isolate number.)  
Select F1 or click “Add spot” to do this.*


8. Select F12 or click validate on right side of screen.

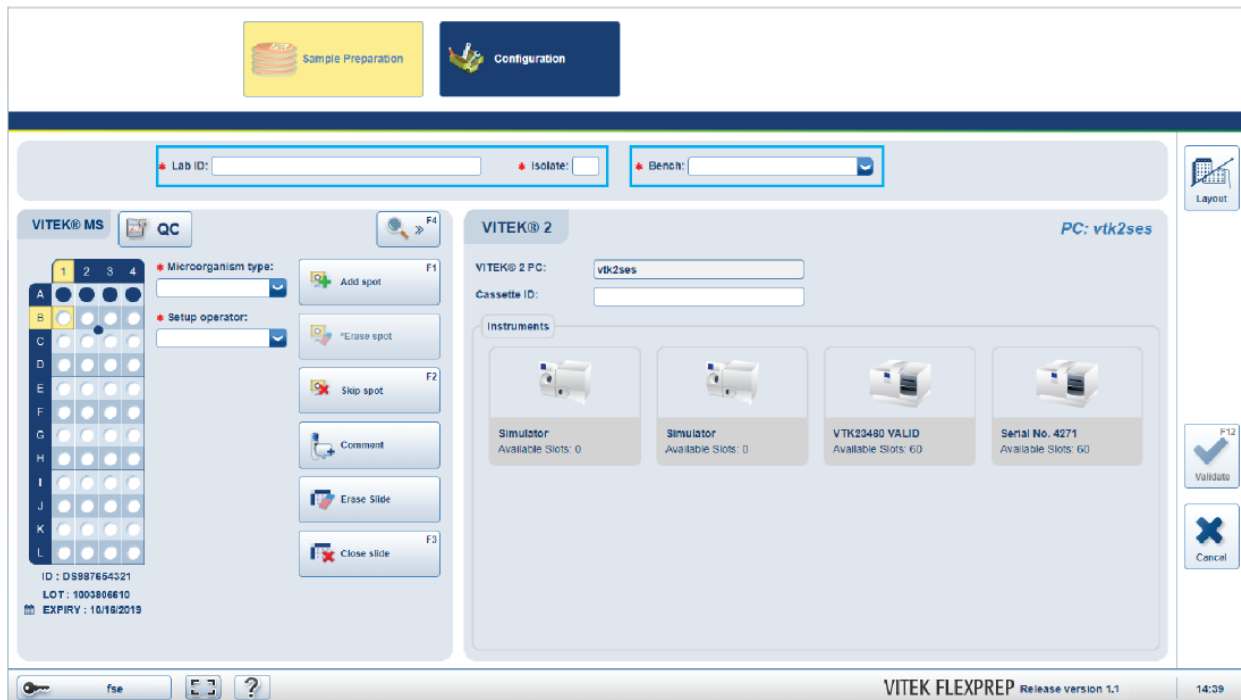
*Note: You may skip a spot if desired, select F2 or click “Skip spot” then “OK”. The acquisition will continue to the next spot on the slide.  
If you need to edit information you must select the spot and click erase spot. You may reenter the information for that spot, then select F12.*







9. Click Close slide or select F3 to send slide.

*Note: \*\*\*You can rescan the slide if you want to change or add anything to it as long as you haven't loaded it yet.\*\*\**

10. Ensure no slides pending review in the instrument before loading (check clipboard).

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
Button	Function
 Add spot F1	Adding a Spot
 *Erase spot	Erasing a Spot
 Skip spot F2	Skipping a Spot
 *Reagents / Comment	Editing Reagent/Slide Information (if Reagent/Slide Traceability is enabled) Adding or Editing Spot Comments
 Erase Slide	Erasing a Slide
 Close slide F3	Closing a Slide

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
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### ***VITEK MS Acquisition Station***

1. Put on a new pair of gloves
2. DOUBLE CLICK Vitek MS Acquisition window
3. Enter User name/password  
**Note:** Check Application Login Information in Microbiology Internal Manual or bench for posting of user name and current password
4. Click Validation Check Mark or press Enter
5. Click Open button on the upper right side of the computer screen
6. Pull out slide adaptor once the Vitek MS door is open
7. Remove the old slide from the adaptor
  - If slide removed has unused acquisition group(s) - place on partial sorting shelf
  - If all acquisitions on slide removed have been used (all spots finished) – discard slide
8. Put the new slides to be tested slide in the position 1,2,3,4
9. Scan the slide id#
10. Load the slide into Vitek MS machine (barcode to the left side) with one hand push down the metal bar and the other hand slide in the dented side into the machine
11. Press Start button to start the acquisition (if unable to select Start option, you may not have sent your slide information from the prep station, you can send it now)
12. Place MS worksheet on clipboard labeled as “Currently running”

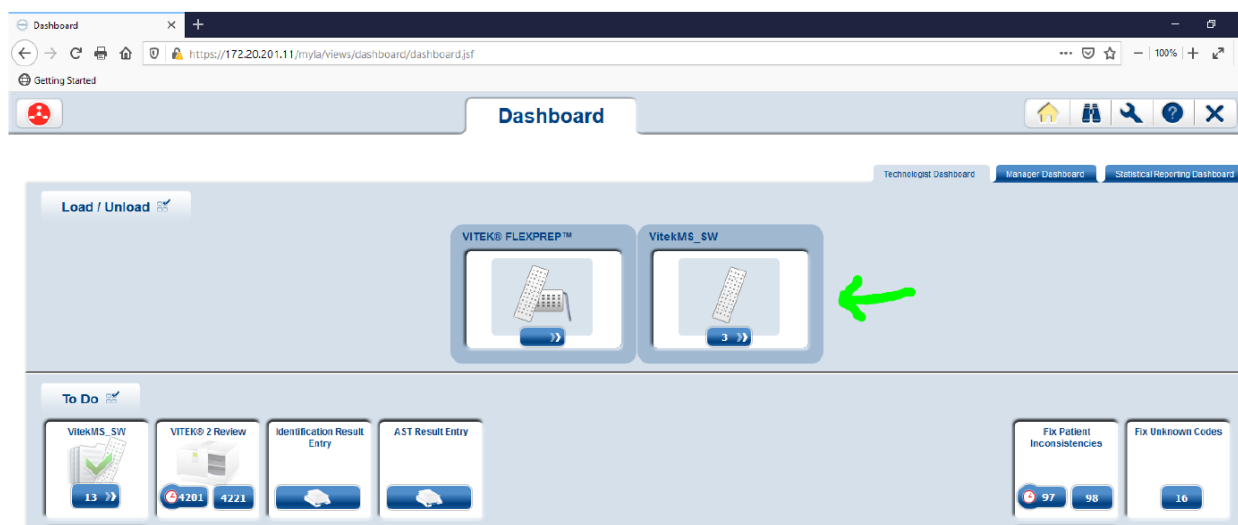
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### **MYLA (Result Review)**

1. Click Dashboard-Window Internet Explorer (MYLA) window
2. Enter User Name
3. Enter Password



**Note:** Check Application Login Information in Microbiology Internal Manual or bench for posting of user name and current password

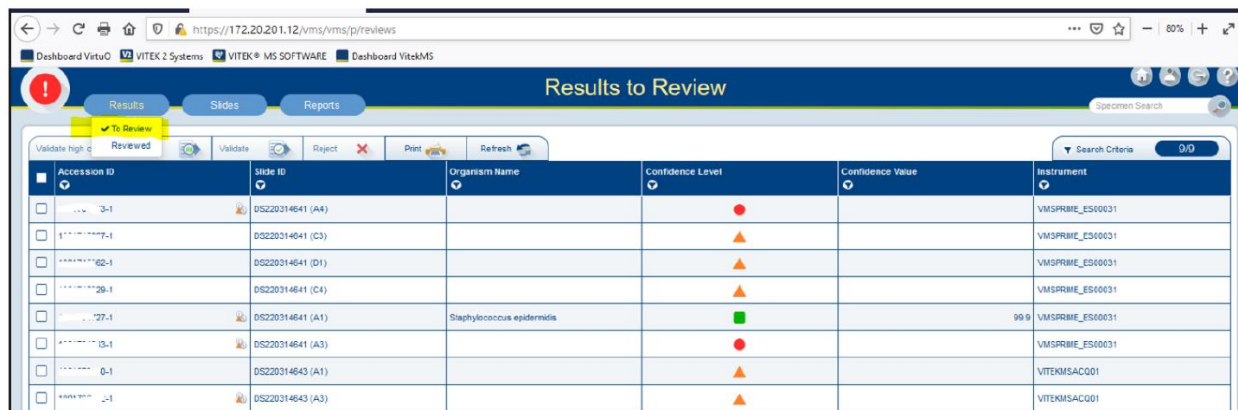
4. Click Login button or press Enter
5. Click Vitek MS Software icon



6. Dashboard window will appear
7. Click on Results and the dropdown menu will give us a selection. Choose “To Review” to review the acquisition
8. VITEK Results to Review will appear




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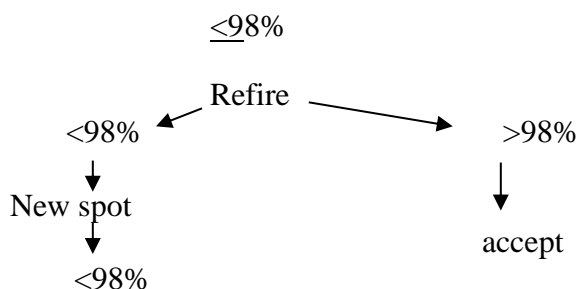


Accession ID	Slide ID	Organism Name	Confidence Level	Confidence Value	Instrument
3-1	05220314641 (A4)		Red dot		VMSPRIME_ES00031
4-1	05220314641 (C3)		Yellow triangle		VMSPRIME_ES00031
5-1	05220314641 (D1)		Yellow triangle		VMSPRIME_ES00031
6-1	05220314641 (C4)		Yellow triangle		VMSPRIME_ES00031
7-1	05220314641 (A1)	Staphylococcus epidermidis	Green square	99.9	VMSPRIME_ES00031
8-1	05220314641 (A3)		Red dot		VMSPRIME_ES00031
9-1	05220314643 (A1)		Yellow triangle		VITEKMSAC001
10-1	05220314643 (A3)		Yellow triangle		VITEKMSAC001

- Filter by Setup Bench by selecting the funnel next to Bench.
- Results can also be filtered by Slide ID, Instrument and Review date. Click on the filter icon to add further details for filtering
- Review results according to confidence level
- In Confidence Level field, click the filter icon and select Red dot (low) as 1<sup>st</sup> step
- Circle the lab# [on the worksheet](#) for further re-fire.
- Isolates that also show a red dot on the Maldi Instrument screen cannot be refired, mark as “no identification” and repeat the spot.
- Click the checkmarks next to the specimen you wish to reject and click “Reject x” in the action bar above the results window.
- Select yellow triangle (medium) as 2<sup>nd</sup> step
- View the result by clicking the **Accession ID#**. Print screen showing the possible organisms. Refer to the MS manual for commonly occurring low discrimination results. Where possible, choose the designated organism. Choose one id only by clicking the round circle on the first left row. Click the appropriate box. Return to MS review. Do this for each organism with a yellow triangle. For those with vastly different organism ID options and/ or implausible ID’s, return to MS review and [reject](#). Circle the lab# on the worksheet for the rejected isolates that need to be re-fired

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18. Check the confidence % for all the isolates with green squares. Any that are <98% should be re-fired. Circle the lab# [on the worksheet](#) for refire as 3<sup>rd</sup> step
19. Print all the results by using printer icon
20. If  $\geq 98\%$ , continue to next step.
21. Select all isolates  $\geq 98\%$  or which have been completely reviewed. Click the green check mark (Review Selected Results icon) to complete the review.
22. Refiring: Take MS worksheet to the Acquisition station computer. Log on. Using the mouse, select all of the circles which have been circled and click Start.
23. After re-firing review the results as above. Those that are still red circles, examine the graph to access the quality of the peaks. If the peaks are good, choose other ID method. If the peaks are not good, an ID may not be acquired due to be a poor prep. It may be worthwhile doing another preparation.
24. For those that are yellow triangles, follow through as above
25. Click the Vitek MS Review to go back previous page
26. All the results should be reviewed and cleared from the review list
27. Put the Matrix and formic acid into the fridge at the end of the day. Close the system
28. Note: You can return to the dashboard at any time by clicking on the ‘house-shaped’ icon
29. If looking for a result by specimen, click on the binoculars in MYLA, then choose tab “Specimen” and enter LIS number (can scan barcode) and click on Search. Drag the blue slide down to view.




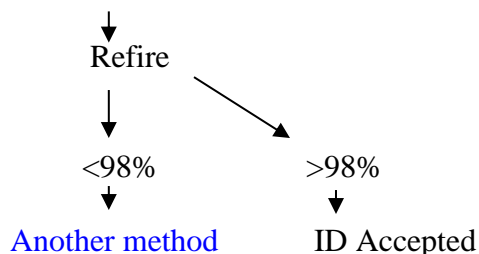
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**Tips:**

- Mucoid colonies: use sterile swab to take off the mucoid layer and then use the loop to pick the colonies.
- Extract all the Yeast with formic acid.
- Make confluent monolayer on the spot

***If results are not transferring, it may be due to demographic inconsistency, see Appendix A for troubleshooting.***


### ***Updating an Isolate Number***

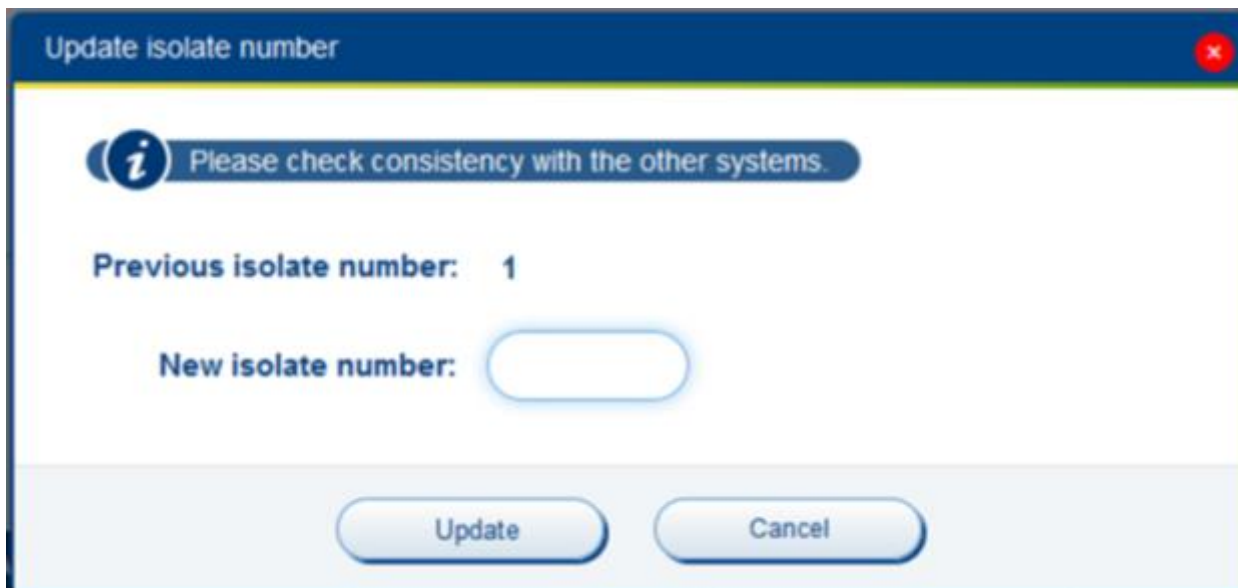
If you used the wrong isolate number when entering your slide information you can change it **before** validating the isolate.

**IMPORTANT:** If you update an isolate number in the VITEK® MS Software, make sure you also update it in VITEK® 2 to ensure consistency between the two systems.

1. In the Specimen Search screen, or from the “To Review” screen, select an Accession ID for which you want to update the isolate number
2. Click Update Isolate Number. The Update Isolate Number window is displayed.  
Note: The Previous isolate number is displayed for information.
3. Enter the New isolate number.

**IMPORTANT:** You cannot choose an isolate number for which a result has already been sent to the bioMérieux middleware.



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4. Click Update. The isolate number is updated. The icon is displayed along with a tooltip indicating the initial isolate number and the new one.

If the updated isolate number was a duplicate, the icon is removed.

Any existing interpretation remains unchanged and is still linked to the same colony. New interpretation results received for the updated isolate are linked to new isolate number (same colony).

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## ***MALDI Organism Identification Acceptance***

### **Bacteria/Yeast**


1. Common isolates with  $\geq 98\%$  report as id by MYLA assuming that the colonial morphology or gram appearance matches the identification
2. Uncommon isolates with MYLA confidence level of  $\geq 98\%$  - report as presumptive with additional in-house testing and confirmation by PHL as necessary assuming that the colonial morphology or gram appearance matches the identification.  
See file “Vitek MS Uncommon bacteria to confirm at PHOL” located in the T:drive  
T:\microbiology\Vitek MS\
3. For identifications that appear as “low discrimination” by MYLA that has a list of a few isolates of the same genera, report as the complex or species based on the full list of Vitek MS organism identifications, Myla organism names and our LIS translated names:  
[VITEK-MS-V2-speciesList Created-Translated list](#)
4. For isolates identified as 50/50 *Klebsiella pneumoniae/variicola*, release identification as *Klebsiella variicola*.
5. Always report the following as presumptive with additional testing for confirmation by PHOL assuming that the colonial morphology or gram appearance matches the identification
  - *Cardiobacterium hominis*
  - *Mycoplasma* species (*Mycoplasma bovis*, *Mycoplasma hyorhithinis*)
  - *Yersinia pestis*
  - *Bacillus anthracis*
6. Never to accept the following organism identifications from MYLA – report Gram stain morphology only
  - *Prevotella oris*
  - *Raoultella ornithinolytica*
  - *Listeria innocua*
  - *Listeria grayii*
7. For any Group A Beta strep, Group C Beta strep and Group G Beta strep identified by MALDI, confirmation by Agglutination is needed before reporting.

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8. For NLF *E coli* identified by MALDI, confirmation by ] is needed before reporting.
- If **Wellcolex Shigella Latex** is negative, report *E.coli*
  - If **Wellcolex Shigella Latex** is positive test with Remel *Shigella* serology and follow reporting algorithm in the table below:

Wellcolex	Remel	Further Workup
POS	POS	<ul style="list-style-type: none"> <li>• Release <i>Shigella species</i> with isolate comment \shcn “presumptive isolated. NOTE: Occasionally, some E. coli may identify as presumptive Shigella species using our current in-lab methodology; confirmation by Public Health Lab reference methodology to follow.”</li> <li>• Send to PHOL to confirm</li> </ul>
POS	Neg	<ul style="list-style-type: none"> <li>• Repeat serology to confirm</li> <li>• Repeatable, set up Vitek GNI</li> </ul> <p><b>Shigella spp ID:</b></p> <ul style="list-style-type: none"> <li>• Release <i>Shigella species</i> with isolate comment \shcn “presumptive isolated. NOTE: Occasionally, some E. coli may identify as presumptive Shigella species using our current in-lab methodology; confirmation by Public Health Lab reference methodology to follow.”</li> <li>• Send to PHOL to confirm</li> </ul> <p><b>NOT a Shigella spp ID:</b></p> <ul style="list-style-type: none"> <li>• Send to PHOL for ID as follow: <ul style="list-style-type: none"> <li>○ Non-stool sites: report GNB ID to follow</li> </ul> </li> </ul> <p><b>Stool specimen: follow Enteric manual</b></p>



9. Any enteric pathogens identified by Vitek-MS require additional testing

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MS Organism Identification	Additional Testing
<i>Salmonella species</i> <i>Salmonella species</i> not <i>S. typhi</i> , not <i>S. paratyphi</i> <i>Salmonella paratyphi</i> A <i>Salmonella typhi</i>	Perform serotyping from BA subculture (same day if possible)
NLF <i>E.coli</i> → possible <i>Shigella species</i>	If <a href="#">Wellcolex Shigella Latex</a> positive, report presumptive ID, then <a href="#">Sub to BA</a> , test with <a href="#">Remel Shigella serology</a> .
NSF <i>E.coli</i> → possible O157	Subculture to BA for: <a href="#">Oxoid E.coli O157 Latex</a> (same day if possible)
<i>Campylobacter C. jejuni</i> or <i>C.coli</i>	Gram stain if atypical colony.
<i>Yersinia</i> <sup>1,2</sup> (any species)	N/A
<i>Vibrio</i> <sup>1,2</sup> (any species)	Gram stain Subculture to BA for Oxidase(+)
<i>Plesiomonas</i> <sup>1,2</sup> or <i>Aeromonas</i> <sup>1,2</sup> (any species)	N/A

<sup>1</sup>Send presumptive identification result when additional testing is completed and send all enteric pathogens to PHL to confirm identification except *C. jejuni* or *C.coli*

<sup>2</sup>If identified by routine specimen processing, proceed to report isolate despite the fact no request for isolation was received.

## 10. Any *Streptococcus pneumoniae* identified by MALDI >98% require confirmation.



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- 99.9% *S.pneumoniae* by MS that are bile soluble or optochin sensitive (only one manual confirmation test required) are confirmed and do not require further work.
- 98-99.8% *S.pneumoniae* ID by MS that are bile insoluble and optochin resistant are sent out as presumptive *S. viridans* and send to PHL for identification confirmation.

#### **If MALDI 99.9% *S.pneumoniae***

Bile solubility	Optochin	Reporting	Send to PHOL
Soluble	Sensitive	<i>S.pneumoniae</i>	No
Soluble	Resistant	presumptive <i>S.pneumoniae</i>	Yes
Insoluble	Sensitive	presumptive <i>S.pneumoniae</i>	Yes
Insoluble	Resistant	presumptive <i>S.pneumoniae</i>	Yes

#### **If MALDI 98-99.8% *S.pneumoniae***

Bile solubility	Optochin	Reporting	Send to PHOL
Soluble	Sensitive	presumptive <i>S.pneumoniae</i>	Yes
Soluble	Resistant	presumptive <i>S.pneumoniae</i>	Yes
Insoluble	Sensitive	presumptive <i>S.pneumoniae</i>	Yes
Insoluble	Resistant	<b>presumptive <i>Viridans strep.</i></b>	Yes

**LIS** References: Q3232098 & Q3063546

11. The following *Neisseria species* specimens must be confirmed by [\[API NH\]](#) before reporting.

- *Neisseria species* - Sterile sites
- *N. gonorrhoeae* - ALL sites
- *N. meningitidis* - ALL sites

If a matching ID is not obtained by API (discrepant result), report isolate as follows:

MS ID	API ID	Site	Reporting if discrepant result	Verify in LIS?
<i>Neisseria species</i>	<i>N.meningitidis</i>	Sterile sites	<i>Neisseria</i> , NOS further report to follow <a href="#">from PHOL</a>	YES
<i>Neisseria species</i>	<i>N.gonorrhoeae</i>	Sterile sites	<i>Neisseria</i> , NOS further report to follow <a href="#">from PHOL</a>	YES
<i>N. gonorrhoeae</i>	Non- <i>N.gonorrhoeae</i>	All sites	<i>Neisseria</i> , NOS further report to follow <a href="#">from PHOL</a>	YES
<i>N. meningitidis</i>	Non- <i>N.meningitidis</i>	All sites	<i>Neisseria</i> , NOS further report to follow from <a href="#">PHOL</a>	YES


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### *Successful Vitek MS identification Tips*


The following tips will help achieve successful identification for spots made and create and efficient workflow.

1. Less is more: ensure a **uniformly thin** layer is applied for successfully readings.
2. Cover entire spot: having the laser read protein rather than a blank area will shorten the time required reading each spot. Hitting blank (reflective) areas also damages the laser
3. BA is preferable: use a colony from blood agar where possible to spot rather than pigmented colonies (e.g. MAC)
4. Avoid agar: Avoid picking up agar when sampling growth for identification as this can lead to misidentification.
5. Avoid specimen: Avoid picking up the inoculated specimen when sampling growth for identification as this can lead to misidentification.
6. Use film growth for BC: an early film growth may be used for sterile sites such a blood but be careful to avoid picking up blood by sampling the film away from the primary inoculum
7. Do NOT re-fire red spots: for any red spots, do not re-fire. Alternatively, re-spot and consider using formic acid.
8. Consider FA for mucoid colonies: For mucoid colonies not achieving a successful ID, consider using formic acid.

### *Vitek MS Etiquette*

Setting up runs:

- Remember to record the slide # on the MALDI worksheet
- Match the isolate # with the number on the MALDI worksheet
  - Do not use the same isolate number twice once it has already been **transferred**
- Perform smaller batches and more frequent loads

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
- Leave a MALDI worksheet by the Vitek MS if slide is in progress

#### Analyzing runs:

- Review your MALDI results soon after it's completed its run
  - Do not leave for extended periods of time without reviewing in MYLA
  - If you will be gone for an extended period, find a technologist willing to review for you
- Remember to remove your MALDI worksheet when finished with Vitek MS
- **Reject** all unwanted IDs after each run
- Place finished worksheets in the save basket when complete
- Place worksheet for "partially" used MS slides in the "Partial" basket near the machine that was used

#### Finished runs:

- Notify the following bench when your run is complete (re-fires completed)
- Ensure the colonial morphology is described for the isolate put into the Vitek MS
  - T:test → J: Morph → pick
- For transferred isolates that are not being reported, switch to as corresponding letter (1=A, 2=B..) and **verify isolate**.
- Ensure the MALDI ID correspond/match the morphology/gram stain of the sample

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## Vitek MS Quality Control

Each Vitek MS requires a positive and negative control QC performed daily for each microorganism type. One bacterial and one yeast isolate will be used for positive quality control. A spot with matrix alone will be used for the negative control.

**This QC shall also be performed after every fine-tuning or when any maintenance is done.**

The following quality control strains are to be used on each day of testing with the expected results below. Results will be documented in Total QC.

VITEK MS DAILY QUALITY CONTROL	
QC ORGANISMS	EXPECTED RESULTS
<i>Enterobacter aerogenes</i> 13048	<i>Enterobacter aerogenes</i> <sup>2</sup>
<i>Candida glabrata</i> MYA-2950	<i>Candida glabrata</i> <sup>2</sup>
No organisms (matrix only)	No ID <sup>1</sup>

<sup>1</sup> If the negative control gives an identification result, visually check the surface of the MS slide to ensure it is clean and repeat negative control

<sup>2</sup> Repeated low confidence identification following proper procedures and technique may indicate calibration may be required.

If unexpected results are achieved follow [Out of Range Results procedure](#) in the Quality Control Manual.

**The daily VITEK MS1 QC is assigned to WASPLab bench technologist and VITEK MS2 QC is assigned to MISC bench technologist. The daily QC is set up on the first slide to be loaded to each MS equipment. The benches responsible for daily QC should repeat and/or follow up with any QC failure, as well as the proper documentation of daily QC result in Total QC software.**


## Vitek MS Maintenance

Routine maintenance is required as scheduled below. All results are documented in the [TQC](#).

### Weekly Maintenance

#### **Desiccant Check**

Check colour of desiccant is orange. If desiccant is changing colour to pink, change the desiccant.

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### ***Area Cleaning***

Clean dusk from Vitek MS area including acquisition station and monitor.

### ***Vitek MS Fine-tuning***

When fine-tuning is being performed ensure to place signs on BOTH Vitek MS Machines.

For machine undergoing Fine-tuned post with sign:

MONITORING OF THIS INSTRUMENT WILL BE PERFORMED FIRST THING  
THIS MORNING  
PLEASE LOAD SLIDES IN MACHINE #1 / #2, UNTIL FURTHER NOTICE

For the machine not undergoing Fine-tuning post with sign:

THE OTHER MALDI MACHINE IS BEING MONITORED THIS MORNING.  
PLEASE ENSURE:  
BLOOD CULTURES & INFECTION CONTROL HAVE FIRST PRIORITY FOR  
LOADING

The calibration status of all instruments are continuously monitored by technical support. A technical support representative will contact the laboratory when an instrument is drifting and fine tuning is required. Fine tuning can be requested by the lab when identification performance is sub-optimal e.g. numerous red spots, extremely slow or multiple failed controls

To perform fine-tuning


- 1) Prepare **two** complete acquisition groups (32 spots plus the control spots) of a new Vitek MS slide from a fresh *E. coli* ATCC 8739 plate.
- 2) Scan the slide and enter “Ecoli8739” into each of the 32 positions on the slide at the Prep Station
- 3) Load the slide into the instrument to be fine-tuned and close, notify the technical support person that it has been loaded
- 4) Do NOT start the run
- 5) Technical support will fine tune the instrument remotely and will contact us when it is complete and ready for use

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- 6) Fine-tuning date must be documented in [TQC](#)

### Monthly Maintenance

#### ***Adapter Cleaning***

Clean adapter with kimwipe.


#### ***O ring and Seal Check***

Open the door and check the condition of both the O-ring and sealing face of the front door. Wipe away moderate soiling with a clean kimwipe. If badly contaminated, use an alcohol to help clean the door seal.




### Yearly Maintenance

Yearly Maintenance is performed by Vitek MS technical staff. A PM visit must be arranged annually.

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### Related Documents:

Document name	Document code
<a href="#">Vitek MS Worksheet</a>	
<a href="#">Vitek MS Worksheet with Controls</a>	
<a href="#">VITEK-MS-V2-speciesList Created-Translated list</a>	Policy: #MI\MSID
<a href="#">International Journal of Systematic and Evolutionary Microbiology</a> (For Official/approved organism name changes)	
<a href="https://lpsn.dsmz.de/">https://lpsn.dsmz.de/</a> (User friendly website to review name changes)	

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## Record of Edited Revisions

### Manual Section Name: Vitek MS MALDI-TOF Manual


Page Number / Item	Date of Revision	Signature of Approval
Updated workflow	November 04, 2013	Dr. T. Mazzulli
Updated MALDI acceptance guide	November 19, 2013	Dr. T. Mazzulli
Monitoring and Fine tuning procedure added	December 16, 2013	Dr. T. Mazzulli
If $\geq 98\%$ , write the result on the MS worksheet. Put a checkmark if 99.9%. Update some process	December 16, 2013	Dr. T. Mazzulli
Annual Review Date	March 31, 2014	Dr. T. Mazzulli
Fix hyperlink in TOC: MALDI Organism Identification Acceptance. Modified header	June 06, 2014	Dr. T. Mazzulli
Removed duplicate steps 31- 63 from Myla Added Vitek MS etiquette section To Vitek MS monitoring and fine tuning added “Monitoring” paragraph at top of section, and “Note” with 4 bullets	June 10, 2015	Dr. T. Mazzulli
-Updated monitoring procedure to verification procedure. -#3 verification: verification slide to be put at 8am #4 added link to email to hotline for verification/finetuning -Removed testing with strep isolate if maldi has an inordinate amount of unidentified isolates. Added point #10 hotline team reply via email	July 16, 2015	Dr. T. Mazzulli
Beginning of Section: <b>Vitek MS Verification and Fine-tuning</b> added instructions for posting signage on both Vitek MS Machines.	September 17, 2015	Dr. T. Mazzulli
Under Organism acceptance added: For Beta hemolytic strep: GAS, GCS, GGS identified by MALDI, confirmation by <a href="#">Streptococcus Latex Agglutination</a> is needed before reporting.	October 01, 2015	Dr. T. Mazzulli
Under verifications: Added separate email templates for Vitek MS 1 and Vitek MS 2	October 7, 2015	Dr. T. Mazzulli
MS organism ID acceptance added #7 criteria for <i>Streptococcus pneumoniae</i>	November 30, 2015	Dr. T. Mazzulli
Added to organism ID list: always send for confirmation	February 22, 2016	Dr. T. Mazzulli

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as per Biomerieux recall Feb 22 <sup>nd</sup> 2016 for: <i>Yersinia pestis, Bacillus anthracis</i>		
Added to Maldi Etiquette: <ul style="list-style-type: none"> <li>Place finished worksheets in the save basket when complete</li> <li>Place worksheet for “partially” used MS slides in the “Partial” basket near the machine that was used</li> </ul>	April 12, 2016	Dr. T. Mazzulli
Removed monitoring values for indication of fine tuning. Document verification and fine tuning in SoftMic weekly. Added Maintenance section with weekly and monthly and yearly maintenance required.	May 16, 2016	Dr. T. Mazzulli
Annual Review Added Related Documents table with link to Vitek myla ID table and MS worksheet	June 10, 2016	Dr. T. Mazzulli
Removed Total Peaks Value communicated and documentation needed. (due to Myla upgrade, no longer required)	December 6, 2016	Dr. T. Mazzulli
Maldi verification spots increased from 16 to 32 spots with controls.	March 13, 2017	Dr. T. Mazzulli
Updated requirements on confirming ID of <i>S.pneumoniae</i> to include performing optochin.	April 6, 2017	Dr. T. Mazzulli
Annual Review	June 10, 2017	Dr. T. Mazzulli
Addition of daily QC section including yeast control.	July 4, 2017	Dr. T. Mazzulli
Addition of expanded Neisseria acceptance rules based on species and sites. Added enteric media additional testing Expanded procedure for NLF <i>E.coli</i> .	August 15, 2017	Dr. T. Mazzulli
Annual Review Updated link to location verification email to technical support. Added section :Tips for successful identification	April 26, 2018	Dr. T. Mazzulli
Annual Review	April 23, 2019	Dr. T. Mazzulli
Addition of MALDI mold procedure and QC organism	January 06, 2020	Dr. T. Mazzulli
Safety Precautions added to VITEK MS Slide spot preparation & VITEK MS Slide spot preparation for	February 21, 2020	Dr. T. Mazzulli


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mold cultures		
Annual Review	May 10, 2020	Dr. T. Mazzulli

**Full document review included in all updates. Biennial review conducted when no revision had been made within 2 years.**


Page Number / Item	Date of Revision	Edited by:
Minor formatting change	April 11, 2021	Jessica Bourke
Added note for kleb pneumo/variicola 50/50	May 18, 2021	Wayne Chiu
References for ID name changes added	June 23, 2021	Jessica Bourke
Clarified S. pneumo confirmation chart	July 6, 2021	Wayne Chiu
Removed username/password	Aug 18, 2021	Wayne Chiu
Included Prevotella oris to list of “never report from MALDI”	Sep 17, 2021	Wayne Chiu
Included Raoultella ornithinolytica to list of “never report from MALDI”	Dec 24, 2021	Wayne Chiu
Removed username/password information from VITEK MS Prep Station, VITEK MS Acquisition Station and MYLA applications	February 25, 2022	Oliver Li
Removed Appendix A, move to BC manual	June 6, 2022	Wayne Chiu
Removed Haemophilus haemolyticus from limitation list	November 1, 2022	Wayne Chiu
Updated MALDI spot preparation instructions	November 23, 2022	
Updated send to PHOL for MALDI ID S. pneumoniae		
Updated SOP to reflect changes to Software Update to VITEK MS v3.2 in sections: VITEK MS Prep Station, MYLA (Result Review)	June 8, 2023	Qin Liu
Added section: Updating an Isolate Number		
Minor formatting	July 21, 2023	Qin Liu
Added/Updated tips for Successful Vitek MS identification (page 19):		
<ul style="list-style-type: none"> <li>• Avoid agar</li> <li>• Avoid specimen</li> <li>• Updated the tip for use film growth for BC : avoid</li> </ul>	November 1, 2023	Qin Liu

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Page Number / Item	Date of Revision	Edited by:
picking up blood by sampling the film away from the primary inoculum.		
Added VITEK MS daily QC duties on page 21 Resolved the inconsistency in handling finished worksheets	October 08, 2024	Oliver Li
Deleted Verification procedure	February 06, 2025	Oliver Li
Added Vitek MS QC after every fine-tuning and any maintenance done and to document in TQC Changed word from biannual to biennial	June 5, 2025	Karin Schoer

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