

Contract No:Co2403079

Date:09/03/2024

Letter of Authorization

Manufacturer: Atlas Medical GmbH Ludwig-Erhard-Ring 3, 15827Blankenfelde-Mahlow, Germany Tel: +49 33 70 83 55 030 Email: <u>amug@atlas-medical.com</u>

Regulatory Office: William James House, Cowley Road, Cambridge, CB4 0WX, UK Tel: +44 1223 858 910 Fax: +44 1223 858 524 Email: <u>info@atlas-site.co.uk</u>

Middle East Site: Sahab Free Zone Area P. O. Box 204, Amman 11512, Jordan. Tel.: +962 6 4026468 Fax: +962 6 4022588 Email: <u>info@atlas-medical.com</u>

Agent: San Medico Republic of Moldova, city Chisina +37368228890

Atlas Medical, hereby appoint the above mentioned agent to import, register and distribute Atlas Medical Products in Maldova

Appointment Conditions:

- 1. This appointment is valid for 3 year from the above mentioned date.
- 2. Either Party can cancel this appointment by giving the other party a 60 day notice.

On behalf of the Manufacturer General Manager Haya Amawi

Atlas Medical Quality Diagnostic Products

Atlas Medical: Ludwig-Erhard-Ring 3, 15827 Blankenfelde-Mahlow, Germany. Tel: +49 33 70 83 55 030 Regulatory Office: William James House, Cowley Road, Cambridge, CB4 0WX, UK. Tel: +44 1223 858 910 Middle East Site : King Abdullah the Second Industrial Estate, Street 19, Sahab Free Zone Area, P.O. Box: 204, Amman 11512, Jordan



GMED certifie que le système de management de la qualité développé par

GMED certifies that the quality management system developed by

ATLAS MEDICAL GmbH Ludwig-Erhard-Ring 3 15827 Blankenfelde-Mahlow GERMANY

pour les activités for the activities

Conception et développement, fabrication et vente de dispositifs médicaux de diagnostic in vitro .

Design and Development, Manufacturing and Sales of in vitro diagnostic medical devices.

réalisées sur le(s) site(s) de performed on the location(s) of

Voir addendum

See addendum

est conforme aux exigences des normes internationales complies with the requirements of the international standards

ISO 13485: 2016

Début de validité / Effective date October 9th, 2023 (included) Valable jusqu'au / Expiry date : October 8th, 2026 (included) Etabli le / Issued on : October 9th, 2023



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GMED N° 36655–2 Ce certificat est délivré selon les règles de certification

Ce certificat est délivré selon les règles de certification GMED / This certificate is issued according to the rules of GMED certification

e sur Renouvelle le certificat 36655-1

GMED • Société par Actions Simplifiée au capital de 300 000 € • Organisme Notifié/Notified Body n° 0459 Siège social : 1, rue Gaston Boissier - 75015 Paris • Tél. : 01 40 43 37 00 • gmed.fr





Ce certificat couvre les activités et les sites suivants :

This certificate covers the following activities and sites:

French version :

Conception et développement, fabrication et vente de dispositifs médicaux de diagnostic *in vitro* à usage professionnel et/ ou d'autodiagnostic, dans les domaines du groupage sanguin, de la microbiologie, de la biochimie, de la toxicologie, de l'oncologie, de la cardiologie, de l'histologie, de l'endocrinologie et des maladies infectieuses, dans les techniques d'Agglutination/ ELISA/ Tests rapides/ Colorimétrie/ Disques antibiotiques.

English version:

Design and Development, Manufacturing and Sales of in vitro diagnostic medical devices for professional use and/or for selftesting, in the field of Immunohematology, Microbiology, Biochemistry, Toxicology, Oncology, Cardiology, Histology, Endocrinology Biosensors and Infectious diseases, in techniques of Agglutination/ ELISA/ Rapid tests/ Colorimetry/Antibiotic disks.

ATLAS MEDICAL GmbH Ludwig-Erhard-Ring 3 15827 Blankenfelde-Mahlow GERMANY

French version: **Siège social, responsable de la mise sur le marché** *English version: Headquarter, legal manufacturer*

Sahab Industrial Zone Area King Abdullah II Industrial City Amman 11512 JORDAN

French version: **Conception, fabrication et contrôle final** *English version: Design, manufacture and final control*

DocuSigned by

On behalf of the President Béatrice LYS Technical Director



Declaration Ref No: DC21-0035

CE Declaration of Conformity

According to Annex III of the IVD Directive 98/79/EC

We,

Atlas Medical

Head office: Ludwig-Erhard-Ring 3 Blankenfelde-Mahlow, Germany. Tel: +49 - 33708 – 3550 30 Email: <u>info@atlas-medical.com</u>

Middle East Site: Sahab Free Zone Area, P. O. Box 212555, Amman, Jordan. Tel.: +962 6 4026468 Fax: +962 6 4022588 Email: <u>info@atlas-medical.com</u>

Declare our responsibility that the following product:

See Attached list

- Comply with all essential requirements (AnnexI) of the IVD Directive 98/79/EC. This compliance has been properly documented and covers the items listed in Annex I of the IVD Directive.
- This product is produced under Atlas quality system (ISO13485:2016) issued by GMED: Certificate N⁰.: 36655 rev 1 Expiry Date: October 8 th.2023
- Comply with the essential requirements of following standards (EN 18113-1, -2,-4:2011, EN ISO 15223:2016, EN ISO 23640:2015, EN ISO 14971:2019, ISO 2859/1:1999, EN ISO 13612:2002, EN ISO 13641:2002.

And Intended for In-Vitro Professional use only.

Manufacturer Atlas Medical Ludwig-Erhard-Ring 3 Blankenfelde-Mahlow, Germany.

Blankenfe	elde-Mahlow, G	Germany.	Atlas Medical	
Atlas	Issue date	Date of review	Quality Diagnostic Products Management approval	MRXDO10F.10
Medical	March.2021	09.03.2021		08.02.2011



CE Declaration of Conformity

According to Annex III of the IVD Directive 98/79/EC

Product Description8.00.02.0.0100 : ASO Latex Kit, 100 Tests (4ml Latex, 2x1.0ml controls).8.00.00.0.0100: CRP Latex Kit, 100 Tests (4ml Latex, 2x1.0 ml Controls)8.00.04.0.0100: RF Latex Kit, 100 Tests (4ml Latex, 2x1.0ml controls)8.00.17.0.0100: D-Dimer Latex Kit, 100 Tests8.00.13.0.0300 : Streptococcus Latex Kit, 6 Groups, 6x50 Tests (5x1.5ml Latex(A,B,C,G,F), 1x3ml Latex(D), 1x1.0ml Positive Control, 1x2ml Extraction Reagent E, 1x1.5ml Extraction Reagent 1, 1x1.5ml Extraction Reagent 2, 2x2.5ml Extraction Reagent 3, Stirring Sticks, Glass Slide).8.00.18.3.0500 : RPR Syphilis (Coarse Grain) Kit, 500 Tests (10 ml latex, 2x1ml control)Without card, stirring sticks.

8.00.18.3.1000 RPR Carbon Antigen (Coarse Grain) Kit, 1000 Tests (Reagent only).

Atlas Medical Quality Diagnostic Products



Date: 15.10.2021

CE Declaration of Conformity

Name and address of Manufacturer	Atlas Medical GmbH	
	Ludwig-Erhard-Ring 3, 15827 Blankenefelde-Mahlow	
	Germany .	
	Tel: +49(0)33708355030	
	Email: info@atlas-medical.com	

Atlas Medical GmbH declared our his own responsibility that the following IVD medical devices:

Product Code	Product Name	CMDN	
8.17.003.0300 Atlas Periodic Acid Schiff (PAS) Stain Kit, 3x100ml		GMDN code	
8.17.004.0300	Atlas Iron Stain Kit, 3x100ml	43587	
8.17.009.1000	Atlas Gram Stain Kit	43587	
8.17.010.0750	Atlas ZN (Kinyoun) stain pack , 3x250ml	43733	
8.15.144.0250	Atlas ZN Decolouriser, 250 ml /Bottle	43587	
8.17.015.0500	Atlas Diff-3 Stain.	43587	
8.17.016.1000	Atlas Papanicolau Stain Pack.	43587	
8.17.110.0250	Atlas Papanicolau Stain EA35, 250 ml /Bottle.	43587	
8.17.111.0250	Atlas Papanicolau Stain EA36, 250 ml /Bottle	43587	
8.17.112.0250	Atlas Papanicolau Stain EA65, 250 ml /Bottle.	43587	
8.17.114.0250	Atlas Papanicolau Stain EA50, 250 ml /Bottle.	43587	
8.17.115.0250	Atlas Papanicolau Stain OG6, 250 ml /Bottle.	43587	
	Atlas Reticulocytes stain (Methylene Blue) , 1000 ml	43587	
8.17.014.1000	/Bottle	43587	
8.15.037.0250	Atlas Eosin Y (1%) Stain, 250 ml/Bottle	42507	
8.15.038.0250	Atlas Eosin Y (5%) Stain, 250 ml/Bottle.	43587	
8.15.041.0250	Atlas Field Stain (Solution A), 250ml/Bottle	43587	
8.15.042.0250	Atlas Field Stain (Solution B), 250ml/Bottle	43587	
8.15.043.0750	Atlas Field Stain Kit 3x250ml (250ml Fixing Reagent ,	43587	
	250ml Eosin Reagent, 250ml Methylene Blue Reagent).	43587	
8.15.047.0250	Atlas Giemsa Stain, 250 ml/Bottle.	12527	
8.15.059.0250	Atlas Haematoxylin Harris Stain , 250 ml/Bottle	43587	
3.15.069.0250	Atlas Leishman Stain , 250 ml/Bottle.	43587	
3.15.069.1000	Atlas Leishman Stain , 1000 ml/Bottle.	43587	
3.15.074.0250	Atlas Lugol's Iodine, 250 ml/Bottle.	43587	
.15.078.0250	Atlas May Grunwald Stain, 250 ml/Bottle.	43587	
	Atlas New Methylene Blue for Reticulocytes, 250	43587	
3.15.105.0250	ml/Bottle.	43587	
3.15.143.0250	Atlas Wright's Stain, 250 ml/Bottle.		
8.15.146.0100	Atlas Immersion oil, 100 Bottle/Box	43587	
	100 BOLLE/BOX	43587	



Declaration Ref No: DC21-0249

Date: 15.10.2021

Meets the essential requirments of In Vitro Diagnostic Medical Devices Directive 98/79/EC Annex I And

EN ISO 13485 :2016, EN 18113-1, -2,:2011, EN ISO 15223:2016 EN ISO 14971:2019, EN ISO 23640:2015, ISO 2859/1:1999, EN ISO 13612:2002, EN ISO 13641:2002, EN ISO 62366-1+A1:2020.

	Directive 98/79, Other IVDs (Non-annex II, non-self-test).	
Conformity Assesment Route	Directive 98/79/EC , Annex III.	
Name , Address and Identification number of notified body	N/A	

Date of issuance:	15. October.2021	
Place	Atlas Medical GmbH	
Signed by:	Amani AL-Habahbeh	
Position :	Ame	
	Regulatory Affairs Manager	

Atlas Medical Gmbn Ludwig - Erhard Ring 3 15827 Blankenfelde - Mahlow Tel. (0049) 33708 - 355030



STREPTOCOCCAL GROUPING SLIDE

TEST

IVD For *In-Vitro* diagnostic and professional use only

^{8°C} Store at 2° to 8° C

CE

INTENDED USE:

2°C /

ATLAS Streptococcus Latex Kit is used for qualitative detection and identification of the Lancefield group of Streptococci. Reagents are provided for groups A, B, C, D, F and G.

INTRODUCTION

ATLAS Streptococcal test uses an enzyme extraction procedure to release Carbohydrate antigen from Streptococcal cell walls. The antigens are detected using specific antibodies to groups A, B, C, D, F and G Lancefield. These antibodies are coated on latex particles. When the antigen extract is mixed with the latex reagent, agglutination will occur. The agglutination appears as a visible clumping and can be seen macroscopically.

PRINCIPLE

Some well isolated colonies are mixed with chemical extraction reagents to liberate the group antigen. This antigen is spread on different circles of the testing glass slide.

Then latex sensitized with antibodies specific for each group, is added. If the correspondent antigen is present in the sample, the antigen-antibody reaction will cause a visible agglutination (clumping). If a sample shows negative reaction with latex of groups A, B, C, F, and G, select other colonies morphologically similar to the proceeding and threat them with the reagent for enzymatic extraction. Test the obtained antigen with latex for group D. A polyvalent extract of streptococci of the abovementioned groups is supplied as a control for the reliability of the latex reagents.

MATERIALS

MATERIALS PROVIDED

- Extracting Reagent 2: Acetic acid solution, ready to use.
- Extracting Reagent 3: Ammonium carbonate solution, ready to use. Contains sodium azide 0.9 g/L as preservative.
- Extracting Reagent E: lyophilized lisozyme in Tris buffer pH 8.2 + 0.2. Contains non-reactive stabilizer and sodium azide 0.9 g/L as preservative. Before use, dissolve with 2.0 mL of sterile distilled water.
- Latex A: sensitized with antibodies (from rabbit) to streptococci of group A. Ready to use. Contains sodium azide 0.9 g/L as preservative.
- Latex B: sensitized with antibodies (from rabbit) to

streptococci of group B. Ready to use. Contains sodium azide 0.9 g/L as preservative.

- Latex C: sensitized with antibodies (from rabbit) to streptococci of group C. Ready to use. Contains sodium azide 0.9 g/L as preservative.
- Latex D: sensitized with antibodies (from rabbit) to streptococci of group D. Ready to use. Contains sodium azide 0.9 g/L as preservative.
- Latex F: sensitized with antibodies (from rabbit) to streptococci of group F. Ready to use. Contains sodium azide 0.9 g/L as preservative.
- Latex G: sensitized with antibodies (from rabbit) to streptococci of group G. Ready to use. Contains sodium azide 0.9 g/L as preservative.
- Positive Control: Lyophilized. Streptococci antigens of groups A, B, C, D, F and G in physiological saline. Contains non-reactive stabilizer and sodium azide 0.9 g/L as preservative. Before use, dissolve with 1.0 mL of sterile distilled water.
- Test slide.
- Stirring Sticks.
- Package Insert.

NOTE: This package insert is also used for individually packed reagent.

MATERIALS NEEDED BUT NOT PROVIDED

- Water bath.
- Test tube.
- Pipettes.
- Sterile loop.

PACKAGING CONTENT

REF 8.00.13.0.0300 (5x1.5 mL Latex (A, B, C, G, F) ,1x3.0 mL Latex (D), 1x1.0 mL Positive Control, 1x 1.5 mL Extraction Reagent 1 , 1x1.5 mL Extraction Reagent 2 , 2x2.5 mL Extraction Reagent 3, 1x2 mL Extraction Reagent E, Glass Slide, plastic stirring sticks).

STORAGE CONDITIONS

- The reagents should be stored refrigerated between 2 8°C avoiding direct light.
- Never Freeze or expose to elevated temperature.
- The reagent is stable until the expiry date stated on the product label. Do not use the reagents past the expiry date.

PRECAUTIONS

- 1. The reagents are intended *for in vitro diagnostic and professional* use only.
- 2. Do not pipette by mouth.
- 3. Always ensure an acceptable performance of the kit by performing the test on the Positive controls before using the kit.

- 4. Wear protective clothing and disposable gloves when dealing with samples and reagents. Wash hands after operations.
- 5. Test materials and samples should be discarded properly in a biohazard container.
- 6. Wash hands and the test table top with water and soap once the testing is done.
- 7. Test specimens may contain pathogenic organisms and must be handled with appropriate precautions.
- 8. When used in accordance with the principles of Good Laboratory Practice, good standards of occupational hygiene and the instructions in these Instructions for Use, the reagents supplied are not considered to present a hazard to health.
- 9. Do not use the kit if the kit label is not available or damaged.
- 10. Don't use the kit if damaged or the vials are leaking and discard the contents immediately.
- 11. The test should be performed at room temperature in a well let area with very good visibility.
- 12. Do not use the reagent if it contains particles as this may indicate reagent deterioration or contamination.
- 13. The Latex Suspensions and Positive Control contain 0.9g/l sodium azide. Azides can react with copper and lead used in some plumbing systems to form explosive salts. The quantities used in this kit are small; nevertheless when disposing of azide-containing materials they should be flushed away with large volumes of water.
- 14. In accordance with the principles of Good Laboratory Practice it is strongly recommended that extracts at any stage of testing should be treated as potentially infectious and handled with all necessary precautions.
- 15. Extraction Reagents 2 and 3 contain a weak acid and a mild irritant respectively. Avoid direct contact by wearing suitable protective equipment. If the material comes into contact with the skin, mucous membranes or eyes immediately wash the area by rinsing with plenty of water.

REAGENT PREPARITION

Latex reagents and extracting reagents 1, 2, and 3 and are ready to use. Bring the reagents to room temperature before use, shake the latex reagents gently to obtain a homogenous suspension of particles. After opening, the reagents are stable until the expiry date if kept as indicated in "STORAGE CONDITIONS". Extracting Reagent E and Positive control are lyophilized and must be re-suspended in sterile distilled water before use. If stored at 2-8 ° C and preserved from contamination, reagents are stable for 3 months.

SPECIMEN AND SAMPLE PREPARATION

For a correct identification it is important that the colonies (which must be well isolated on blood agar) are picked up fresh.

Before serological analysis, it is advisable to observe the hemolytic activity and set up a slide with Gram stain to ensure the purity of the strain to be tested.

PROCEDURES

Allow all reagents and samples to reach room temperature (18-30°C) before use.

- A. Technique with Chemical Extraction
- 1. Transfer **30 μL (one drop) of Extracting Reagent 1** into a labelled test tube.
- Pick up 5-6 colonies with a stirring stick, being careful not to pick up part of the culture medium. Add colonies into the test tube and mix to obtain a homogeneous suspension.
- 3. Transfer 30 µL (one drop) of Extracting Reagent 2.
- Let stand for at least 5 minutes at room temperature. Do not exceed 10 minutes. A prolonged extraction time decreases the sensitivity of the test.
- 5. Transfer **60 μL (two drops) of Extracting Reagent 3** and mix. Use within 15 minutes.
- 6. Re-suspend the latex reagent to be used (i.e. A, B, C, F, and/or G) by shaking the vial.
- 7. Holding the dropper vertically, add 1 free-falling drop of latex in one circle of the glass slide. Repeat this operation for each latex to be used.
- 8. Transfer 15 µL of antigenic extract in each circle.
- 9. Using a clean stirring stick, mix and spread the reaction mixture carefully. Discard the used stirring stick.
- Tilt and rotate the glass slide. After one minute, observe each circle for evidence of agglutination (clumping). Later agglutinations should be considered as nonspecific.
 NOTE: If all results are negative, proceed with the technique for identification of Group D Streptococci.

B. Direct Technique

(This procedure is able to identify about 70% of Group D strains).

- 1. Transfer **30** µL (a drop) of Extracting Reagent **3** in a circle of the slide.
- 2. Pick up 2-3 colonies with a clean stirring stick, being careful not to pick up part of the culture medium, and carefully mix them in the same circle of the slide.
- 3. Add a drop of Latex D.
- Tilt the slide for 1 minute. At the end observe each circle for the presence or absence of agglutination. Later agglutinations should be considered as nonspecific.
 NOTE: If negative results are obtained continues with enzymatic extraction technique.

C. Technique with Enzymatic Extraction

(This procedure is able to identify more than 95% of group D strains)

 Distribute, after reconstitution, 60 μL (two drops) of Extracting Reagent E into a labelled test tube.

- 2. Pick up 2-3 colonies with a clean stirring stick, being careful not to pick up part of the culture medium. Insert colonies into the test tube and mix to obtain a homogeneous suspension.
- 3. Incubate at 37° C for 10 minutes.
- Holding the dropper vertically, add 1 free-falling drop of Latex D in one circle of the glass slide.
- 5. Add **15 µL of antigenic extract** in one circle.
- 6. Using a clean stirring stick, mix and spread the reaction mixture carefully. Discard the used stirring stick.
- 7. Tilt and rotate the glass slide. After one minute, observe each circle for evidence of agglutination (clumping). Later agglutinations should be considered as nonspecific.

Quality Control

Use the positive control and saline as if they were extracted from a sample. The absence of reactions (respectively positive or negative) is index of alteration of the reagents and / or controls .

READING THE RESULT

A. Technique with Chemical Extraction

Positive: If Agglutination appears in the test circle with latex A, B, C, F or G respectively.

Negative: Fine particles appear in the test circle with latex A, B, C, F or G respectively with **no agglutination or clumping**.

B. Direct Technique

Positive: If Agglutination appears in the test circle with latex D. **Negative: Fine particles** appear in the test circle with latex D with **no agglutination or clumping**.

C. Technique with Enzymatic Extraction

Positive: If Agglutination appears in the test circle with latex D. **Negative: Fine particles** appear in the test circle with latex D with **no agglutination or clumping**.

NOTE: An insufficient amount of bacterial culture used can cause false negative results.

PERFORMANCE CHARACTERISTICS

Sensitivity

The identification with chemical extraction technique of groups A, B, C, F and G streptococci, performed both on lyophilized collection strains and on clinical isolations, has showed a sensitivity of 98%.

The identification of group D with direct technique has showed a sensitivity of 74.3%.

The identification of group D with enzymatic extraction has showed a sensitivity of 92%.

REFERENCES

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PPI1415A01

Rev G (18.10.2023)

REF	Catalogue Number	ľ	Temperature limit
IVD	In Vitro diagnostic medical device	\wedge	Caution
¥	Contains sufficient for <n> tests and Relative size</n>	<u> </u>	Consult instructions for use (IFU)
LOT	Batch code		Manufacturer
I	Fragile, handle with care		Use-by date
	Manufacturer fax number		Do not use if package is damaged
3	Manufacturer telephone number	R	Date of Manufacture
*	Keep away from sunlight	ſ	Keep dry
CONTROL +	Positive control	Control -	Negative control



GRAM STAIN PACK

IVD For in -vitro diagnostic and professional use only

Store at Room Temperature

INTENDED USE

Gram Stain used for differentiate between gram positive and gramnegative bacteria.

INTRODUCTION

Gram staining is used to differentiate bacterial species into two large groups (Gram-positive and Gram-negative) based on the physical properties of their cell walls.

PRINCIPLE

Gram-positive bacteria have a thick mesh-like cell wall made of peptidoglycan (50-90% of cell wall), which stains Blue while gramnegative bacteria have a thinner layer (10% of cell wall), which stains pink. Gram-negative bacteria also have an additional outer membrane which contains lipids, and is separated from the cell wall by the periplasmic space. There are four basic steps of the Gram stain, which include applying a primary stain (crystal violet) to a heat-fixed smear of a bacterial culture, followed by the addition of a trapping agent (Gram's iodine), rapid decolorization with alcohol or acetone, and *counterstaining* with safranin or basic fuchsin.

Crystal violet (CV) dissociates in aqueous solutions into CV⁺ and chloride (Cl⁻) ions. These ions penetrate through the cell wall and cell membrane of both gram-positive and gram-negative cells. The CV⁺ ion interacts with negatively charged components of bacterial cells and stains the cells Blue.

lodine (I $^-$ or I_3 $^-$) interacts with CV+ and forms large complexes of crystal violet and iodine (CV–I) within the inner and outer layers of the cell. Iodine is often referred to as a mordant, but is a trapping agent that prevents the removal of the CV-I complex and therefore color from the cell.

When a decolorizer such as alcohol or acetone is added, it interacts with the lipids of the cell membrane. A gram-negative cell will lose its outer membrane and the lipopolysaccharide layer is left exposed. The

CV–I complexes are washed from the gram-negative cell along with the outer membrane. In contrast, a gram-positive cell becomes dehydrated from an ethanol treatment. The large CV–I complexes become trapped within the gram-positive cell due to the multilayered nature of its peptidoglycan. The decolorization step is critical and must be timed correctly; the crystal violet stain will be removed from both gram-positive and negative cells if the decolorizing agent is left on too long (a matter of seconds).

After decolorization, the gram-positive cell remains Blue. and the gram-negative cell loses its Blue. color. Counterstain, which is usually positively charged safranin or basic fuchsin, is applied last to give decolorized gram-negative bacteria a pink or red color.

MATERIALS

MATERIALS PROVIDED Crystal Violet

- Crystal Violet.
- Gram lodine.
- Gram Decolouriser.
- Counterstain Safranin O.

Note: This package insert is also used for individually packed reagent.

Packaging Content

- REF
 8.17.009.0400
 (1x100ml
 Crystal
 Violet,
 1x100ml
 Iodine

 Solution,
 1x100ml
 Decolouriser,
 1x100ml
 Safranin O)
- REF 8.17.008.1000 (1x250ml Crystal Violet, 1x250ml Iodine Solution, 1x250ml Decolouriser, 1x250ml Safranin O)
- REF 8.17.009.1000 (1x250ml Crystal Violet, 1x250ml Iodine Solution, 1x250ml Decolouriser, 1x250ml Safranin O)

- REF 8.15.032.0250 (1x250ml Crystal Violet)
- REF 8.15.049.0250 (1x250ml lodine Solution)
- REF 8.15.051.0250 (1x250ml Decolouriser) REF 8.15.125.0250 (1x250ml Safranin O)

STORAGE AND STABILITY

- Store at room temperature.
- Stain Solution is stable up to the printed expiry date.
- Keep the bottles tightly closed to prevent air oxidation.

PRECAUTIONS

- The reagent may cause eye, skin and respiratory tract irritation; so protective clothing should be worn when handling this reagent.
- The reagent is intended for in vitro diagnostic use only.
- Do not use this reagent if the label is not available or damaged.
- Test materials and samples should be discarded properly in biohazards container.
- This reagent is considered toxic, so do not drink or eat beside it.
- Wash hands and test table top with water and soap once the testing is done.

PROCEDURE

- 1. immerse the heat fixed smears with Crystal Violet and allow to stain for up to 1 minute.
- 2. Wash with tap water.
- 3. Flood the smear with Gram Iodine for 2 minutes.
- 4. Wash with tap water.
- 5. Decolorize the smear for few second only.
- 6. Wash thoroughly with tap water.
- 7. Counterstain with Safranin O for up to 2 minutes.
- 8. Wash and allow to dry.
- 9. Examine under microscope using oil immersion objective

RESULTS

- Gram positive organisms (Blue).
 - Gram negative organisms (Red).

ATLAS Medical GmbH Ludwig-Erhard Ring 3 15827 Blankenfelde-Mahlow Germany Tel: +49 - 33708 – 3550 30

Email: Info@atlas-medical.com Website: www.atlas-medical.com

PPI2112A01

Rev C (27.03.2023)

