

BIO-DIFF 3 REAGENT

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IVD In vitro diagnostic medical device

Classified acc. to Regulation (EU) 2017/746 - Class A device

Blue component of Bio-Diff kit INSTRUCTIONS FOR USE

BASIC UDI number	385889212HPC3010302HMCA						
EMDN code	W0103010302	V0103010302					
REF Catalogue number	Volume	UDI-DI number	REF Catalogue number	Volume	UDI-DI number		
BD3-0T-100	100 mL	03858888824419	BD3-OT-1L	1000 mL	03858888824433		
BD3-0T-500	500 mL	03858888824426	BD3-0T-2.5L	2500 mL	03858892120057		



Intended use and test principle

Bio-Diff 3 reagent is a component of BioGnost's Bio-Diff kit. Contains azure dyes in phosphate buffer. Other reagents found in the kit is Bio-Diff 1 reagent (fixative), Bio-Diff 2 reagent (red component), and pH 6.8 and 7.2 buffer tablets. Bio-Diff 1 reagent contains methyl alcohol, while Bio-Diff 2 contains eosin Y dye in phosphate buffer. Red and blue components' interaction creates a well known Romanowsky effect that cannot be achieved if each component is being used individually. Purple color indicates the effect's presence. BioGnost's Bio-Diff kit efficiently stains hematology and cytology smears in a short period of time and provides precise staining results, such as results of the May-Gruenwald-Giemsa method. Each component of kit is stabilized and prepared according to the highest standards.

Product description

• BIO-DIFF 3 REAGENT - azure dyes solution in phosphate buffer

Using Bio-Diff 3 reagent as a component of Bio-Diff kit

Additional reagents and materials that can be used in staining:

- BioGnost's immersion oils, such as BioGnost's Immersion oil, Immersion oils types C, A, FF and Tropical Grade
- · VitroGnost slides and coverslips for use in histopathology and cytology
- Other components of Bio-Diff kit: Bio-Diff 1 reagent (catalog number BD1-OT-100, BD1-OT-500, BD1-OT-1L, BD1-2.5L) and Bio-Diff 2 reagent (catalog number BD2-OT-100, BD2-OT-500, BD2-OT-1L, BD2-OT-2.5L) and Buffer tablets pH 7.2 (catalog number PT-72-50, PT-72-100) or Buffer tablets pH 6.8 (catalog number PT-68-50, PT-68-100)

Preparation of solutions

Buffer solution, pH 6.8 or 7.2

Dissolve 1 pH 6.8 buffer tablet in 1 liter of distilled water while stirring. Filter the solution.

Note

Pour the reagents into glass staining jars (type Coplin, Hellendahl or Schifferdecker) and return them to the original bottles after staining. Close well.

Blood smear/bone marrow sample staining procedure

1.	Let the smear dry	
	Note: Prepare the peripheral blood smear by draining blood from a fresh blood sample	
2.	Dip the smear into Bio-Diff 1 reagent	5 x 1 second
3.	Decant the excessive reagent from the smear onto filter paper	
4.	Dip the smear into Bio-Diff 2 reagent	3 x 1 second
	Note: extend the incubation period if a stronger hue of red/purple is required	do 5 x 1 second
5.	Decant the excessive reagent from the smear onto filter paper	
6.	Dip the smear into Bio-Diff 3 reagent	6 x 1 second
	Note: decrease the incubation period if a stronger hue of red/purple is required	5 x 1 second
7.	Rinse the smear in pH 6.8 buffer solution	1 min (with agitation)
8.	Dry the smear	

Staining method for parasitology (Leishmania, Toxoplasma, Microsporadia) and microbiology samples (Cryptosporidium, Pneumocystis carinii)

1.	Dip the smear into Bio-Diff 1 reagent	1 min
2.	Decant the excessive reagent from the smear onto filter paper	
3.	Dip the smear into Bio-Diff 2 reagent	25 seconds
4.	Decant the excessive reagent from the smear onto filter paper	
5.	Dip the smear into Bio-Diff 3 reagent	25 seconds
6.	Rinse the smear in pH 7.2 buffer solution	1 min (with agitation)
7.	Dry the smear	

Sperm staining procedure

Preparing the sperm smear: Add 15 μ L of fresh sperm sample on one side of the glass slide and create a thin and homogeneous smear. Let the smear dry (at least 10 minutes).

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1	. Dip the smear into Bio-Diff 1 reagent	5 x 1 second
2	2. Decant the excessive reagent from the smear onto filter paper	
3	3. Dip the smear into Bio-Diff 2 reagent	5 x 1 second
4	Decant the excessive reagent from the smear onto filter paper	
	i. Dip the smear into Bio-Diff 3 reagent	5 x 1 second
- 6	6. Rinse the smear in pH 7.2 buffer solution	1 min (with agitation)
7	. Dry the smear	

In order to create a permanent sample, apply appropriate type of DPX medium on both stained and dried smear (BioMount DPX medium for covering/mounting cover slides). Cover the smear with VitroGnost cover glass.

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Head - homogeneous dark purple Acrosome - light purple Mid piece and tail - dark purple Background - light pink

Histological sections staining procedure

a) preparation of histological sections for staining

- Fix (Formaldehyde NB 4%, Formaldehyde NB 10%) and process the tissue sample
- Embed the tissue in a paraffin block (BioWax 52/54, BioWax 56/58, BioWax Plus 56/58, BioWax Blue)
- Cut the paraffin block into 4-6 μ m thin slices and mount on a VitroGnost microscope slide

b) sample staining procedure

Note: do not use Bio-Diff 1 reagent (used as fixative for non-histology samples)

1.	Deparaffinize the section in xylene (BioClear) or in a xylene substitute (BioClear New)	3 exchanges, 10 min each
2.	Rehydrate in 100% alcohol (Histanol 100)	2 exchanges, 5 and 3 min
3.	Rehydrate in 95% alcohol (Histanol 95)	2 min
4.	Rehydrate in distilled/demineralized water	2 min
5.	Dip the section into Bio-Diff 2 reagent and gently stir	7 seconds
6.	Dip the section into Bio-Diff 3 reagent and gently stir	5 seconds
7.	Rinse in Buffer solution pH 7.2	1 min (with agitation)
8.	Decant the excessive reagent from the section onto filter paper	
9.	Dehydrate and differentiate in 95% alcohol (Histanol 95) while gently stirring	10 seconds
10.	Dehydrate in 100% alcohol (Histanol 100)	1 min
11.	Clear in xylene (BioClear) or xylene substitute (BioClear New)	2 exchanges, 5 min each

Immediately after clearing, apply an appropriate BioMount covering/mounting medium. If BioClear xylene was used, use one of BioGnost's xylene-based mountants (BioMount. BioMount High, BioMount M, BioMount DPX, BioMount C, or universal BioMount New). If BioClear New xylene substitute was used, the appropriate mountant is BioMount New. Cover the section with a VitroGnost cover glass.

Cytobacteriology samples staining procedure (urine, punctates, CSF)

1.	Let the cytology smear dry	
2.	Dip the smear into Bio-Diff 1 reagent	5 seconds
	Note: Incubate CSF for a longer period of time	1 min
3.	Decant the excessive reagent from the smear onto filter paper	
4.	Dip the smear into Bio-Diff 2 reagent	3 x 1 seconds (CSF 2 x 1 second)
	Note: extend the incubation period if a stronger hue of red/purple is required	do 5 x 1 second
5.	Decant the excessive reagent from the smear onto filter paper	
6.	Dip the smear into Bio-Diff 3 reagent	6 x 1 seconds (CSF 2 x 1 second)
	Note: decrease the incubation period if a stronger hue of red/purple is required	5 x 1 second
7.	Rinse the smear in pH 7.2 buffer solution	1 min (with agitation)
8.	Dry the smear	

Results for staining of cytological and histological samples

Nuclei - red to purple

Lymphocytes - plasma is colored blue Monocytes - plasma is colored grey-blue Neutrophil granulocytes - light purple Eosinophil granulocytes - bright red to red-brown Basophil granulocytes - dark purple to black

Thrombocytes - purple Erythrocytes - reddish

Blood parasites - red (nuclei), blue (cytoplasm)

Helicobacter pylori – dark blue Tissue cellular elements - blue to pink

Limitations

This product is intended for professional laboratory use for diagnostic purposes only. Deviations from the staining procedure described in this Instruction for use may cause differences in staining results.

Sample preparation and diagnostics

Use only appropriate instruments for collecting and preparing the samples. Process the samples using modern technology and mark them clearly. Be sure to follow the manufacturer's handling instructions. To avoid errors, staining, mounting of the slides, and diagnosis can only be carried out by qualified personnel. Use a microscope equipped according to medical diagnostic laboratory standards.

If a serious incident occurs during use of this product or as a result of its use, please report it to the manufacturer or authorized representative and competent authority.

Safety at work and environmental protection

Handle the product in accordance with occupational health and environmental protection guidelines. Used and expired solutions must be disposed of as special waste following national guidelines. Reagents used in this procedure can pose a danger to human health. The examined tissue samples are potentially infectious, and it is necessary to take the measures needed to protect human health in accordance with the guidelines of good laboratory practice. It is mandatory to read and act according to the information and warning signs printed on the product label and in the Safety Data Sheet, which is available on request.

Storage, stability, and shelf life

Upon receipt, store the product in a dry place and well-closed original packaging at a temperature of +15 °C to +25 °C. Do not freeze or expose to direct sunlight. After first opening, the product can be used until the specified expiry date, if stored properly. The expiration date is printed on the product label.

Literature

- 1. Beck, R.C. (1938): Laboratory Manual of Heamtological Technique, Philadelphia, W.B. Saunders & Co.
- Dacie, J. et Lewis S. (1995): Practical haematology, 4th ed., London, Churchill Livingstone.
- Giemsa, G. (1922): Das Wesen der Giemsa-Farbung, Zentralb f Bakt; 89, p 99-106.
- Kieman, J.A. (2008): Histological and histochemical methods: Theory and Practice, 4th ed., Bloxham, Scion Publishing Ltd.
- May, R. et Grünwald L. (1909): Über die Farbung von Feutchpraparaten mit meiner Azur-Eosine methode, Deutsche med Xschr, 35, p 1751-1752.

Warnings and precautions regarding the materials contained in the product: Not a dangerous substance or mixture according to Regulation (EC) No. 1272/2008.

BD3-X-IFU, EN2, 10 November, 2025, ISP							
***	Manufacturer	LOT	Batch code	Ţį	Consult instructions for use	C€	European conformity
W	Date of manufacture	REF	Catalogue number	<u> </u>	Caution	UDI	Unique device identifier
\Box	Use-by date	°C-¶-°C	Temperature limit	IVD	In vitro diagnostic medical device		

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Version	Description / reason for change	Date
2.	Addition of: the QR code, table with the Warnings and precautions, revised table with the symbols	10.11.2025.