

Stay healthy, live better

PRIMA Lab SA is a **modern Swiss Company** developing simple and precise CE certified diagnostic in vitro medical devices, for a **rapid check of people's health** and **their pets**.

PRIMA also offers **rapid tests for health care professionals**, providing point of care analysis to improve their services.

Prevention has never been so simple, handy and accessible to everybody and everywhere: **in just 5-10 minutes** you can get an accurate answer about your health status and that of your pet.

Stay healthy, live better!





REF	DESCRIPTION	N° OF TESTS	SAMPLE	SHELF LIFE
100077-1	CELIAC TEST	1 TEST	WHOLE BLOOD	24 MONTHS
800017-1	HELICOBACTER PYLORI TEST	1 TEST	WHOLE BLOOD	30 MONTHS
800016-1	HELICOBACTER PYLORI ANTIGEN TEST	1 TEST	FAECES	30 MONTHS
800040-1	BOWEL FOB TEST	1 TEST	FAECES	30 MONTHS
800062-1	HIV 1/2 SELF-TEST	1 TEST	WHOLE BLOOD	24 MONTHS
800034-1	C-REACTIVE PROTEIN TEST	1 TEST	WHOLE BLOOD	36 MONTHS
800060-1	STREP A TEST	1 TEST	THROAT SWAB	30 MONTHS
100013-5	VAGINAL pH TEST	5 TESTS	VAGINAL SWAB	24 MONTHS
100058-3	URINARY TRACT INFECTIONS TEST	3 TESTS	URINE	24 MONTHS
800080-1	PROSTATE PSA TEST	1 TEST	WHOLE BLOOD	30 MONTHS
820027-1	KIDNEYS ALBUMIN TEST	1 TEST	URINE	30 MONTHS
800025-1	IRON FER TEST	1 TEST	WHOLE BLOOD	30 MONTHS
800018-1	ALLERGY IgE TEST	1 TEST	WHOLE BLOOD	36 MONTHS
100066-1	VITAMIN D TEST	1 TEST	WHOLE BLOOD	24 MONTHS
19806-1	DRUG TEST	1 TEST	URINE	24 MONTHS
820010-x	EARLY PREGNANCY TEST	1-2 TESTS	URINE	30 MONTHS
820050-5	OVULATION LH TEST	5 TESTS	URINE	30 MONTHS
820011-2	MENOPAUSE FSH TEST	2 TESTS	URINE	30 MONTHS
100082-1	SPERM TEST	1 TEST	SPERM	24 MONTHS
100088-5	AMNIOCHECK	5 TESTS	AMNIOTIC FLUID	48 MONTHS
PET100-1	GIARDIA PET TEST	1 TEST	PET FAECES	24 MONTHS

PRIMA® PROFESSIONAL

_	REF	DESCRIPTION	N* OF TESTS	SAMPLE	SHELF LIFE
	100066IP-20P	VITAMIN D RAPID TEST	20 TESTS	WB / S / P	24 MONTHS
	800060IP-20P	STREP A RAPID TEST	20 TESTS	THROAT SWAB	30 MONTHS



REF	DESCRIPTION	N* OF TESTS	SAMPLE	SHELF LIFE
800087-5	KI SCREENING TEST	5 TESTS	DOG'S URINE	18 MONTHS





CELIAC TEST

Self-test to check for the presence of anti-deamidated gliadin (anti-DGP) antibodies, IgA and IgG, in the blood, for celiac disease screening



CELIAC DISEASE

Celiac Disease (CD), also known as "celiac sprue", is a chronic inflammatory disorder of the small intestine, produced by the ingestion of dietary gluten products in susceptible people. It is a multifactorial disease, including genetic and environmental factors. Environmental trigger is represented by gluten while the genetic predisposition has been identified in the major histocompatibility complex region which cause the presence of specific antibodies against gluten and its components. Celiac disease is not a rare disorder like previously thought, with a global prevalence around 1%. The reason of its under recognition is mainly referable to the fact that about half of affected people do not have the classic gastrointestinal symptoms, but they present nonspecific manifestations of nutritional deficiency or have no symptoms at all.

GLUTEN, GLIADIN AND DGP

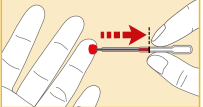
Gluten is a protein found in wheat, composed by glutenin and gliadin. It helps foods maintain their shape, acting as a glue that holds food together. Gliadin is a peptide water-soluble component of gluten and it can pass through the intestinal epithelium. Gliadin is the part of gluten responsible of Celiac Disease. DGP: Deamidated gliadin peptides are the result of the action of Transglutaminase Enzyme and its detection could increase sensitivities, specificities and accuracies of celiac disease screening.

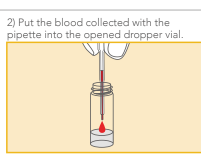
WHO ARE THE INTENDED USERS

Patients with symptoms (abdominal bloating and pain, chronic diarrhea, vomiting, constipation, weight loss, fatigue, irritability), patients with a first degree family member with CD, and people with Type I diabetes.

HOW TO USE IT

1) Take a blood sample after pricking the finger.





3) Add 3 drops of liquid and wait 10 minutes before reading the result.



CONTENT: 1 sealed aluminium pouch containing: 1 test device and 1 desiccant bag; 1 transparent plastic bag containing a pipette for blood collecting; 1 vial with dropper containing the diluent; 2 sterile lancets for blood sampling; 1 alcohol swab and 1 instructions for use leaflet.

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(\land)	VIDEO	REF	DESCRIPTION	NUMBER OF TESTS	SHELF LIFE
TUTORIAL	100077-1	CELIAC TEST	1 TEST	24 MONTHS	

WHY - BENEFITS

Underdiagnosis is common and the quality-of-life impact of symptoms may be severe. Checking the presence of specific antibodies against DGP is a useful tool to detect Celiac disease.

TEST PRINCIPLE

CELIAC TEST is based on an immunochromatographic reaction. If the antibodies (IgA and IgG) are present into the sample, they will react with the deamidated gliadin. A positive result means that the concentration of antibodies is nearby to 20 AU/mL.

TECH SPECS

CUT-OFF	CUT-OFF SENSITIVITY		OVERALL ACCURACY
20 AU/mL	81,25%	94,10%	90,00%

Performance data obtained by two clinical study with 200 participants enrolled. ImmunoCap Thermofisher DGP IgG/IgA (ELISA) assay and Zenit RA Menarini DGP IgG/IgA (Chemiluminescent IA) assay have been utilized as reference methods.

CLINICAL EVIDENCES

1. P.D. Mooney, S.H.Wong, A.J.Johnston, M.Kurien, A.Avgerinos and D.S.Sanders "Increased detection of celiac disease with measurement of Deamidated Gliadin Peptide Antibody befor endoscopy" Cli, Gastroenterology and Hepatology 2015;13:1278-1284 2. N. R. Lewis & B. B. Scott "Meta-analysis: deamidated gliadin peptide antibody and tissue transglutaminase antibody compared as screening tests for coeliac disease" Alimentary Pharmacology & Therapeutics

3. A. Lammi, P. Arikoski, S. Simell, T. Kinnunen, V. Simell, S Paavanen-Huhtala, A. Hinkkanen, R. Veijola, M. Knip, J. Toppari, O. Vaarala, O. Simell, and J. Ilonen "Antibodies to Deamidated Gliadin Peptide in Diagnosis of Celiac Disease in Children" Gastroenterology - JPGN 2015;60: 626–631





HELICOBACTER PYLORI TEST

Self-test for the determination of anti Helicobacter pylori antibodies in whole blood samples



HELICOBACTER PYLORI

Helicobacter pylori (Hp) is a bacterium that chronically infects more than half of the world's population and plays a causative role in the pathogenesis of chronic gastritis, peptic ulcer diseases, gastric cancer and mucosa associated lymphoid tissue lymphoma. The considerable burden of these Hp-related outcomes means that there is an acute demand for accurate diagnosis of this infection. Several detection methods have already been developed, such as culture, histological staining, the urea breath test (UBT) but a simple, non-invasive, inexpensive and accurate diagnostic test remains the goal. The simplest ways to detect infection is checking the presence of specific antibodies (IgG) produced by the immune system after encountering the bacterium and trying to eliminate it. HELICOBACTER PYLORI TEST can detect the presence of these antibodies in few minutes.

WHO ARE THE INTENDED USERS

Everyone who has gastric symptoms; the most common manifestations of these conditions are heartburn or pain in the upper part of the abdomen, especially when the stomach is empty. Other associated symptoms include nausea, feeling full, reflux, loss of appetite, weight loss and diarrhoea. Ulcers can sometimes bleed, causing anemia over long periods of time.

WHY - BENEFITS

In 1994, the National Institutes of Health recognized that most recurrent duodenal and gastric ulcers were caused by H. pylori and antibiotic treatment was recommended. In the same year, the International Agency for Research on Cancer

(IARC) declared H. pylori to be a group I human carcinogen for gastric adenocarcinoma.

TEST PRINCIPLE

HELICOBACTER PYLORI TEST is an immunochromatographic test that detects the presence of anti-Helicobater pylori IgG antibodies in the blood, using special monoclonal antibodies conjugated with gold and integrated into the reactive strip.

TECH SPECS

CUT-OFF	SENSITIVITY	SPECIFICITY	OVERALL ACCURACY
15 AU/mL	95,80%	84,20%	87,00%

Performance data obtained by clinical study with 100 participants enrolled. IBL ELISA kit has been utilized as reference method.

CLINICAL EVIDENCES

1. Mati Moyat, Dominique Velin. Immune responses to Helicobacter pylori infection. World J. Gastroenterol. 2014 may 21; 20(19): 5583-5593IgE." Journal of Allergy and Clinical Immu-

nology. 2014;133(2):589-91

2. Khean-Lee Goh, Wah-Kheong Chan, Seiji Shiota and Yoshio

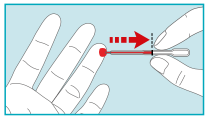
Yamaoka. Epidemiology of Helicobacter pylori Infection and Public Health Implications. Helicobacter, 2011 Sep;16(0 1):1-9.

3. Kyichi Adachi, Tomoko Mishiro, Shino Tanaka and Yoshikazu Kinoshita. Analysis of negative result in serum

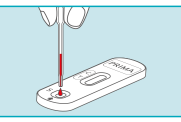
anti-pyllori IgG antibody test in cases with gastric mucosal atrophy. J. Clin Biochem Nutr.2016 Sep;59(2):145-148.

HOW TO USE IT

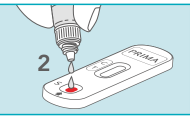
1) Take a blood sample after pricking the finger.



2) Deposit the sample into the specimen well of the cassette.



3) Add 2 drops into the well and wait 5 minutes before reading the result.



CONTENT: 1 sealed aluminium pouch containing: 1 test device and 1 desiccant bag; 1 transparent plastic bag containing a pipette for blood collecting; 1 vial with dropper containing the diluent; 2 sterile lancets for blood sampling; 1 alcohol swab and 1 instructions for use leaflet.

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(\land)	VIDEO	REF	DESCRIPTION	NUMBER OF TESTS	SHELF LIFE
C	TUTORIAL	800017-1	HELICOBACTER PYLORI TEST	1 TEST	30 MONTHS





HELICOBACTER PYLORI ANTIGEN TEST

Self-test for detecting the presence of the Helicobacter pylori antigen in the faeces



HELICOBACTER PYLORI

Helicobacter pylori (Hp) is a bacterium that chronically infects more than half of the world's population and plays a causative role in the pathogenesis of chronic gastritis, peptic ulcer diseases, gastric cancer and mucosa associated lymphoid tissue lymphoma. The considerable burden of these H. pylori-related outcomes means that there is an acute demand for accurate diagnosis of this infection. Several detection methods have already been developed, such as culture, histological staining, the urea breath test (UBT) but a simple, non-invasive, inexpensive and accurate diagnostic test remains the goal. One of the simplest ways to detect infection is by checking the presence of specific antigens for the bacterium that are excreted through the faeces. HELICOBACTER PYLORI ANTIGEN TEST can detect the presence of these antigens in few minutes.

WHO ARE THE INTENDED USERS

Everyone who has gastric symptoms, ulcers can sometimes bleed, causing anemia over long periods of time.

WHY - BENEFITS

In 1994, the National Institutes of Health recognized that most recurrent duodenal and gastric ulcers were caused by *H. pylori* and antibiotic treatment was recommended. In the same year, the International Agency for Research on Cancer (IARC) declared H. pylori to be a group I human carcinogen for gastric adenocarcinoma.

TEST PRINCIPLE

HELICOBACTER PYLORI ANTIGEN TEST is an immunochromatographic test that detects the presence of specific *Helicobacter pylori* antigens in stools by using special monoclonal antibodies conjugated with gold and integrated into the reactive strip.

TECH SPECS

CUT-OFF	SENSITIVITY	SPECIFICITY	OVERALL ACCURACY
10 ng/mL	95,20%	96,60%	96,00%

Performance data obtained by clinical study with 100 participants enrolled. ImmunoCard STAT!® HpSA® has been utilized as reference method.

CLINICAL EVIDENCES

1. Shimoyama T. "Stool antigen tests for the management of Helicobacter pylori infection." World Journal of Gastroenterology. 2013;19(45):8188-91.

 Ana Isabel Lopes, Filipa F Vale, Mónica Oleastro. "Helicobacter pylori infection - recent developments in diagnosis." World Journal of Gastroenterology. 2014 July 28; 20(28): 9299-9313.
 Andreas Mentis, Philippe Lehours, and Francis Francis Megraud.

 Andreas Mentis, Philippe Lehours, and Francis Francis Megraud. "Epidemiology and Diagnosis of Helicobacter pylori infection". Helicobacter. 2015 Sep;20 Suppl 1:1-7.
 Zagari RM, Rabitti S, Eusebi LH, Bazzoli F. "Treatment of Helico-

4. Zagari RM, Rabitti Ś, Eusebi LH, Bazzoli F. "Treatment of Helicobacter pylori infection: A clinical practice update." Eur J Clin Invest. 2018 Jan;48(1). doi: 10.1111/eci.12857

HOW TO USE IT

1) Attach the two adhesive surfaces of the stool-collecting sheet on the toilet seat surface. 2) Dip the stick into the faeces in three different points of the sample.





3) Screw on the cap, immersing the stick in the stool extraction liquid, and shake the vial.



4) Break the end of the cap, freeing the dropper portion. Dispense 3 drops of the diluted stool sample in the well shown on the cassette.



CONTENT: 1 sealed aluminium pouch containing: 1 test device and 1 desiccant bag; 1 vial with collection stick and dripper, containing the diluent; 1 small sheet for the collection of the sample; 1 instructions for use leaflet.







BOWEL FOB TEST

Self-test for the detection of occult blood in the faeces



TECH SPECS

CUT-OFF	SENSITIVITY	SPECIFICITY	OVERALL ACCURACY
40 ng/mL	78,70%	99,00%	92,60%

Performance data obtained by clinical study with 148 participants enrolled. Hb Fecale - Mascia Brunelli kit has been utilized as reference method.

CLINICAL EVIDENCES

1. Van Rossum Leo G. et al.: "Random Comparison of Guaiac and Immunochemical Fecal Occult Blood Tests for Colorectal Cancer in a Screening Population" Gastroenterology, Volume 135, Issue 1, 82 – 90.

Cancer in a Screening ropulation Gastroenteroug, reterne 135, Issue 1, 82 – 90. 2. Faivre J., Dancourt V., Denis B., Dorval E., Piette C., Perrin P., Bidan J.M., Jard C., Jung S., Levillain R., Viguier J., Bretagne J.F.: "Comparison between a guaiac and three immunochemical faecal occult blood tests in screening for colorectal cancer". European Journal of Cancer. 2012; 48(16):2969-76. 3. Benton S.C., Seaman H.E., Halloran S.P.: "Faecal Occult Blood Testing for Colorectal Cancer Screening: the past or the Future". Current Gastroenterology Reports. 2015; 17(2):428 4. European Commission. European guidelines for quality assurance in colorectal cancer screening and diagnosis. First edition 2010

FOB (FAECAL OCCULT BLOOD)

Faecal Occult Blood (FOB) is the presence in the faeces of blood that is not clearly visible to the naked eye. Occult blood may be the consequence of gastro-intestinal disorders such as ulcers, polyps, colitis, diverticulitis, rhagades, haemorrhoids or cancer. The signs of these lesions are often silent in the first phases, so searching for occult blood in the stool is an important screening test, which allows to identify the presence of pathologies in the gastro-intestinal system at an early stage.

WHO ARE THE INTENDED USERS

Everyone, especially over 45 years.

WHY - BENEFITS

Sometimes, the presence of low quantity of blood in stools is not easily noticeable. In these cases BOWEL FOB TEST is useful to detect the presence of blood in the stools even if they look normal in order to prevent more serious diseases.

TEST PRINCIPLE

BOWEL FOB TEST is an immunochromatographic test that detects the smallest amounts of blood in stool samples by using highly specific monoclonal antibodies. These properties mean that the BOWEL FOB TEST does not require particular dietary restrictions before being carried out.

HOW TO USE IT

1) Attach the two adhesive surfaces of the stool-collecting sheet on the toilet seat surface. 2) Dip the stick into the faeces in three different points of the sample.



3) Screw on the cap, immersing the stick in the stool extraction liquid, and shake the vial.



4) Break the end of the cap, freeing the dropper portion. Dispense 3 drops of the diluted stool sample in the well shown on the cassette.



CONTENT: 1 sealed aluminium pouch containing: 1 test device and 1 desiccant bag; 1 vial with collection stick and dripper, containing the diluent; 1 small sheet for the collection of the stool sample; 1 instructions for use leaflet.







HIV 1/2 SELF-TEST

Rapid self-test for the qualitative detection of antibodies against human immunodeficiency virus 1 and 2 (HIV1/HIV2) in whole blood



HIV VIRUS

Human immunodeficiency virus, or HIV, is a pathogen that attacks and suppresses the immune system by specifically affecting white blood cells. Without specific treatment, the virus weakens the subject's immune system to the point of inducing the development of acquired immunodeficiency syndrome, or AIDS. It is a syndrome that can occur in people with HIV even several years after the infection, when the immune system's efficiency dramatically falls, and the body loses its ability to fight even the most common infections.

WHO ARE THE INTENDED USERS

HIV 1/2 SELF-TEST can be used by anyone who has been, or suspects to have been exposed to the virus, people with HIV symptoms or anyone who wants to be aware of a possible infection. Antibody tests can take 23 to 90 days (window period) to detect HIV infection after an exposure.

WHY - BENEFITS

It is very important to support the prevention campaign against HIV, in order to prevent more serious disease, like AIDS, by testing and monitoring the population.

TEST PRINCIPLE

HIV 1/2 SELF-TEST is a rapid immunochromatographic assay, able to specifically detect the presence of antibodies against human immunodeficiency virus 1 and 2 (HIV1/ HIV2) in whole blood sample.

TECH SPECS

SENSITIVITY	SPECIFICITY	OVERALL ACCURACY
99,30%	99,70%	99,60%

CLINICAL EVIDENCES

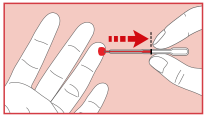
1. https://www.who.int/health-topics/hiv-aids/#tab=tab_1 2. European Centre for Disease Prevention and Control, HIV/ AIDS surveillance in Europe (https://www.ecdc.europa.eu/en/ all-topics-zhiv-infection-andaidssurveillance-and-disease-data/annual-hivaids-surveillance-reports)

3. https://www.hiv.gov/hiv-basics/overview/about-hiv-and-aids/ symptoms-of-hiv

4. Debit et al., "HIV-1 Group O Genotypes and Phenotypes: Relationship to Fitness and Susceptibility to Antiretroviral Drugs" 5. https://www.cdc.gov/hiv/basics/hiv-testing/test-types.html

HOW TO USE IT

1) Take a blood sample after pricking the finger.



2) Put the blood collected with the pipette into the opened dropper vial.



3) Add 3 drops and wait 10 minutes before reading the result.



CONTENT: 1 sealed aluminium pouch containing: 1 test device and 1 desiccant bag; 1 transparent plastic bag containing a pipette for blood collecting; 1 vial with dropper containing the diluent; 2 sterile lancets for blood sampling; 1 alcohol swab and 1 instructions for use leaflet.

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(\land)	VIDEO	REF	DESCRIPTION	NUMBER OF TESTS	SHELF LIFE
E	TUTORIAL	800062-1	HIV 1/2 SELF-TEST	1 TEST	24 MONTHS





C-REACTIVE PROTEIN TEST

Rapid self-test for the semi-quantitative detection of C-Reactive Protein (CRP) in whole blood samples



TECH SPECS

CUT-OFF	SENSITIVITY	SPECIFICITY	OVERALL ACCURACY
10 mg/L	98,70%	96,00%	97,60%

Performance data obtained by clinical study with 127 participants enrolled. Beckam Coulter au680 CRP Latex has been utilized as reference method.

CLINICAL EVIDENCES

1. THOMPSON, D.; MILFORD-WARD, A.; WHICHER, J. T. "The value of acute phase protein measurements in clinical practice". Annals of clinical biochemistry, 1992, 29.2: 123-131. 2. SHAW, A. C. "Serum C-reactive protein and neopterin concentrations in patients with viral or bacterial infection". Journal of clinical

pathology, 1991, 44.7: 596-599 3. C-reactive protein concentrations as a marker of inflammation or infection for interpreting biomarkers of micronutrient status. Vitamin and Mineral Nutrition Information System. Genf: World Health Organization; 2014 (https://www.who.int/nutrition/publications/micronutrients/indicators_c-reactive_protein/en/)

4. https://www.mayoclinic.org/tests-procedures/c-reactive-protein-test/about/pac-20385228

C-REACTIVE PROTEIN

C-Reactive Protein (CRP) is a non specific marker mainly produced by the liver and used to diagnose bacterial diseases and inflammatory disorders. CRP is a very sensitive and fast appearing indicator which could therefore be helpful for deciding an antibiotic treatment.

C-REACTIVE PROTEIN TEST should be performed in case of acute infection symptoms such as feverishness, fever, headaches or weakness.

WHY - BENEFITS

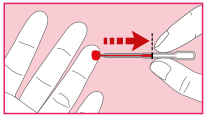
In healthy patients, CRP concentration is lower than 8 mg/L while the concentration level can be higher than 100 mg/L in case of severe infection or during inflammatory process. Intermediate levels, within 8 and 100 mg/L, are concomitant with moåre or less mildly viral or bacterial infections that can be easily overcome by medical treatment. Checking the CRP concentration in blood is useful to verify the presence of possible infections or inflammatory states.

TEST PRINCIPLE

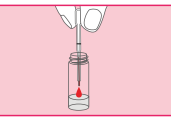
C-REACTIVE PROTEIN TEST is an immunochromatographic assay which detects the CRP level thanks to special monoclonal gold-conjugate antibodies embedded to test strip.

HOW TO USE IT

1) Take a blood sample after pricking the finger.



2) Put the blood collected with the pipette into the opened dropper vial.



3) Add 3 drops and wait 5 minutes before reading the result.



CONTENT: 1 sealed aluminium pouch containing: 1 test device and 1 desiccant bag;, 1 transparent plastic bag containing a pipette for blood collecting; 1 vial with dropper containing the diluent; 2 sterile lancets for blood sampling; 1 alcohol swab and 1 instructions for use leaflet.







STREP A TEST

Self-test for the detection of Group A B-hemolytic Streptococcus in throat swab



STREPTOCOCCUS A

Group A β-hemolytic streptococcus (also known as *Streptococcus pyogenes*) infection of the throat is one of the most common bacterial cause of acute pharyngitis and is also responsible for some skin infections such as impetigo and erysipelas. Most sore throats are caused by viral infections which clear up on their own without antibiotic treatment, which would be useless. While Strep A throat infections sometimes can go away within a few days without treatment, physicians prescribe antibiotics to prevent related complications that can be serious, such as rheumatic fever or acute glomerulonephritis.

WHO ARE THE INTENDED USERS

STREP A TEST can be carried out if there are symptoms such as pain when swallowing, sore throat, red and swollen tonsils, small red patches on the back of the palate, swollen lymph nodes, fever, headache, nausea or vomiting, especially in children.

WHY - BENEFITS

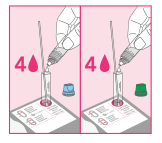
STREP A TEST helps to quickly know whether a sore throat is caused by the bacterium Streptococcus A, which generally requires an antibiotic treatment, or other germs (usually viruses) that do not require this type of treatment.

HOW TO USE IT

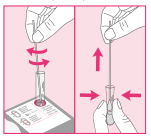
1) Collect a throat sample using the supplied swab and the tongue depressor



2) Put the swab into the extraction tube inserted in the hole on the back of the package and add the two dilution solutions.



3) Mix the solutions and leave the swab inside the tube for 2-5 minutes. Then remove it and squeeze the cotton tip to collect as much liquid as possible.



4) Apply the dropper to the tube and dispense 1 drop into the well.



CONTENT: 1 hermetically sealed protective aluminium pouche containing: 1 test device and a desiccant bag; 1 sterile swab; 1 sterile wooden tongue depressor; 1 empty plastic test tube for sampling, with dropper; 1 vial with dropper containing the STREP A TEST - R1 solution required for 1 test (blue cap); 1 vial with dropper containing the STREP A TEST - R2 solution required for 1 test (green cap) and 1 instructions for use leaflet.

	REF	DESCRIPTION	NUMBER OF TESTS	SHELF LIFE	
U	TUTORIAL	800060-1	STREP A TEST	1 TEST	30 MONTHS

TEST PRINCIPLE

STREP A TEST is an immunochromatographic test that detects the presence of specific streptococcus A antigens in throat swab samples by using special monoclonal antibodies.

TECH SPECS

CUT-OFF	SENSITIVITY	SPECIFICITY	OVERALL ACCURACY
10 ⁴ CFU/mL	91,89%	99,00%	97,90%

Performance data obtained by clinical study with 238 samples tested comparing with another rapid immunochromatographic test used as reference method.

CLINICAL EVIDENCES

1. Banerjee S, Ford C. "Rapid Tests for the Diagnosis of Group A Streptococcal Infection: A Review of Diagnostic Test Accuracy, Clinical Utility, Safety, and Cost-Effectiveness". Ottawa (ON): Canadian Agency for Drugs and Technologies in Health; 2018 May. CADTH Rapid Response Reports.

2. Centers for Disease Control and Prevention. Adult appropriate antibiotic use summary. http://www.cdc.gov/getsmart/ community/ materials-references/print-materials/hcp/adult-approp-summary. pdf 2016.

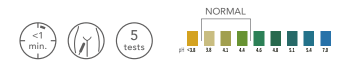
pdf 2016. 3. Llor C., Bjerrum L., Munck A., Cots JM., Hernández S., Moragas A.; HAPPY AUDIT Investigators. "Access to point-of-care tests reduces the prescription of antibiotics among antibiotic-requesting subjects with respiratory tract infections." Respir Care. 2014 Dec.





VAGINAL PH TEST

Rapid self-test for the semi-quantitative determination of pH in female vaginal swab specimens



VAGINAL INFECTIONS AND pH

Vaginal infections are quite common and often a recurring problem among women of all age groups. An indication of abnormal acidity of the vaginal discharge can help evaluate whether the vaginal symptoms are likely caused by an infection that may require follow-up with an healthcare professional. An acidic vaginal pH value of 3.8 to 4.5 is a basic requirement for the optimal functioning of the body system which protects the vagina. This system can effectively avoid colonization by pathogenic germs and the occurrence of vaginal infections.

WHAT IS VAGINAL pH TEST

VAGINAL pH TEST is a rapid dry chemical method for the semiquantitative detection of pH in female vaginal swab specimens to aid in the diagnosis of bacterial vaginosis.

WHO ARE THE INTENDED USERS

Women who feel vaginal discomfort.

WHY - BENEFITS

Vaginitis is a very common feminine disease affecting million of women each year. Vaginitis accounts for over 50% of all gynaecologic office visits. The key to a proper treatment is an early accurate diagnosis to prevent Bacterial Vaginosis complications.

HOW TO USE IT

Insert the head of swab into the vagina and gently press the swab to a point for approximately 10 seconds.

Withdraw the swab from the vagina for testing.

Apply the head of the swab onto the pH area of the Test panel, for at least 5 times.

Read the result immediately while the measurement zone is still wet. Compare the colour in the pH measurement zone with the colour scale on the foil pouch.

TEST PRINCIPLE

A pH paper is used to determine pH levels in specimen obtained through a vaginal swab. As soon as the swab containing the specimen from vagina comes in contact with the pH paper encased in the test panel, a colour change occurs. Different pH levels can result in different grades and shades of colour with each colour grade/shade specifying a particular pH level on the colour scale.

TECH SPECS

RANGE OF MEASUREMENT	SENSITIVITY	SPECIFICITY	OVERALL ACCURACY
3,80 - 7,00	95,00%	99,00%	98,60%

Performance data obtained by clinical study with 220 participants enrolled. PH Vaginal Rapid Test (pH) has been utilized as reference method.

CLINICAL EVIDENCES

1. Pavletic, A. J., Hawes, S. E., Geske, J. A., Bringe, K., & Polack, S. H. (2004). Experience with routine vaginal pH testing in a family practice setting. Infectious Diseases in Obstetrics and Gynecology, 12(2), 63–68.

2. Huppert, J. S., Hesse, E. A., Bernard, M. C., Bates, J. R., Gaydos, C. A., & Kahn, J. A. (2012). Accuracy and Trust of Self-Testing for Bacterial Vaginosis. The Journal of Adolescent Health : Official Publication of the Society for Adolescent Medicine, 51(4), 400–405.



CONTENT: 5 sealed aluminum pouches with printed colour scale each containing 1 test device; 5 sterile vaginal swabs; 1 Instructions for use leaflet.







URINARY TRACT INFECTIONS TEST

Self-test for the detection of leukocytes, blood, nitrites and proteins in urine



TECH SPECS

URINARY TRACT INFECTIONS

Urinary tract infection (UTI) is a collective term that describes any infection involving any part of the urinary tract, namely the kidneys, ureters, bladder and urethra. It is one of the most common infections in local primary care.

WHO ARE THE INTENDED USERS

The test should be performed in presence of the following symptoms: excessive frequency and/or burning during urination, turbid and/or bad-smelling urine.

WHY - BENEFITS

Checking the presence of urinary tract infections is a useful tool to verify the general health status of the urinary system.

TEST PRINCIPLE

URINARY TRACT INFECTIONS TEST detects the presence of non-physiological levels of leukocytes, blood, nitrites and proteins in urine thanks to specifc chemical reactions which are visibile with a colour change of the reactive area.

CUT-OFF				
Leukocytes	9 leu/µL			
Blood: free haemoglobin	0,018 mg/dL or 5 Ery/µL			
Protein	7,50 mg/dL			
Nitrite	0,05 mg/dL			
OVERALL ACCURACY				
Leukocytes	97,60%			
Blood	96,00%			
Protein	88,00%			
Nitrite	100,00%			

Performance data obtained by clinical study with 125 participants enrolled. U120 Urine Analyser has been utilized as reference method.

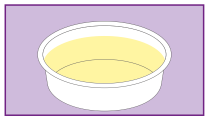
CLINICAL EVIDENCES

1. Chee Wei Tan, MMed, MCFP1and Maciej Piotr Chlebicki, MBBS, ABIM2: "Urinary tract infections in adults" Singapore Med J. 2016 Sep; 57(9): 2. Copp H.L., Schmidt B.: "Work up of Pediatric Urinary Tract Infec-

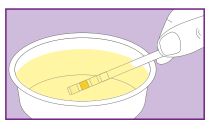
 Copp H.L., Schmidt B.: "Work up of Pediatric Urinary Tract Infection". Urol Clin North Am. 2015 Nov; 42(4): 519–526.

HOW TO USE IT

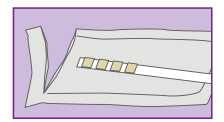
1) The sample is the first urine of the day collected in a clean and dry cup, not provided with the kit.



2) Dip the strip-test for 1-2 seconds.



3) Wait for 2 minutes then check the colours of the strip-test.



CONTENT: 3 sealed aluminium pouches each containing: 1 test strip and 1 desiccant bag; 1 colour chart and 1 instructions for use leaflet.



	REF	DESCRIPTION	NUMBER OF TESTS	SHELF LIFE
AL	100058-3	URINARY TRACT INFECTIONS TEST	3 TESTS	24 MONTHS





PROSTATE PSA TEST

Self-test for the determination of the Prostate-Specific Antigen (PSA) in whole blood samples



PROSTATE SPECIFIC ANTIGEN (PSA)

The Prostate Specific Antigen (PSA) is a protein produced by the epithelial cells of the prostate gland. Its function is to fluidify the ejaculate and to increase sperm mobility. PSA also plays a role in dissolving cervical mucus, allowing sperm to enter the uterus. Small quantities of PSA are normally found in the blood of men with a healthy prostate. PSA levels may increase with prostate pathologies (prostatitis, benign prostatic hyperplasia etc.), naturally with age or after prostate surgery or specific diagnosis tests (such as rectal examination, prostate biopsy etc.).

WHO ARE THE INTENDED USERS

PROSTATE PSA TEST is intended for:

- men who do not have symptoms, aged over 45, who wish to check their prostate;

- men with urinary tract symptoms (difficulty urinating, pain, increased frequency).

WHY - BENEFITS

Checking the level of the PSA is useful to verify prostate health status. In case of cancer, early detection and prevention can be life saving.

TEST PRINCIPLE

PROSTATE PSA TEST is an immunochromatographic assay which detects the PSA thanks to special monoclonal goldconjugate antibodies embedded to test strip.

TECH SPECS

CUT-OFF	SENSITIVITY	SPECIFICITY	OVERALL ACCURACY
4 ng/mL	97,22%	87,13%	92,34%

Performance data obtained by clinical study with 209 participants enrolled. Abbott Alinity has been utilized as reference method.

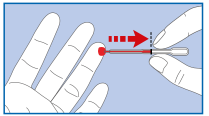
CLINICAL EVIDENCES

1. Catalona W.J., Southwick P.C., Slawin K.M., Partin A.W., Brawer M.K., Flanigan R.C., Patel A., Richie J.P., Walsh P.C., Scardino P.T., Lange P.H., Gasior G.H., Loveland K.G., Bray K.R.: Comparison of percent free PSA, PSA density, and age-specific PSA cutoffs for prostate cancer detection and staging. Urology. 2000 Aug 1;56(2):255-60.

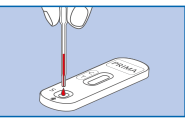
Stitsche H.A., Babaian R.J.: Analytical performance goals for measuring prostate specific antigen. Clin Chem. 1993 Jul;39(7):1525-29.
 Michael J. Barry, M.D: Prostate-Specific-Antigen Testing for Early Diagnosis of Prostate Cancer. The New England Journal of Medicine, 2001; 344:1373-1377.

HOW TO USE IT

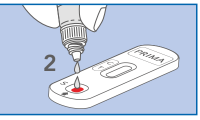
1) Take a blood sample after pricking the finger.



2) Deposit the sample into the specimen well of the cassette.



3) Add 2 drops into the well and wait 5 minutes before reading the result.



CONTENT: 1 sealed aluminium pouch containing: 1 test device and 1 desiccant bag; 1 transparent plastic bag containing a pipette for blood collecting; 1 vial with dropper containing the diluent; 2 sterile lancets for blood sampling; 1 alcohol swab and 1 instructions for use leaflet.

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(\land)	VIDEO	REF	DESCRIPTION	NUMBER OF TESTS	SHELF LIFE
C	/ TUTORIAL	800080-1	PROSTATE PSA TEST	1 TEST	30 MONTHS





KIDNEYS ALBUMIN TEST

Self-test for detecting albumin in the urine



ALBUMIN IN URINE

Albumin is the most abundant plasma protein, formed principally in the liver and constituting up to 2/3 of the 6-8% protein concentration in the plasma. Albumin is responsible for much of the colloidal osmotic pressure of the blood, and thus is very important to regulate the exchange of water between the plasma and the interstitial compartment (i.e. the space between the cells).

The presence of albumin in the urine (albuminuria) may indicate a malfunction of the kidneys and it can accompany kidney disease or heart failure.

WHO ARE THE INTENDED USERS

Everyone who has the suspect of kidney's pathology or with special condition like high blood pressure, diabetes or pregnancy.

WHY - BENEFITS

Checking the presence of albumin protein in urine is useful to verify the general health status.

TEST PRINCIPLE

KIDNEYS ALBUMIN TEST is an immunochromatographic assay which specifically identifies the proteins in human urine and shows if the concentration is equal to or higher than 10 μ g/ml.

TECH SPECS

CUT-OFF	SENSITIVITY	SPECIFICITY	OVERALL ACCURACY
10 µg/mL	98,20%	95,20%	96,90%

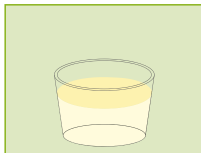
Performance data obtained by clinical study with 99 participants enrolled. Roche Cobas Microalbumin has been utilized as reference method.

CLINICAL EVIDENCES

 Urinary biomarkers of kidney dysfunction. Ann Biol Clin (Paris).
 2015 Mar-Apr;73(2):151-7. doi: 10.1684/abc.2015.1029.
 Lopez-Giacoman, Salvador and Magdalena Madero. "Biomarkers in chronic kidney disease, from kidney function to kidney damage" World journal of nephrology vol. 4,1 (2015): 57-73.
 Sacks, David B et al. "Guidelines and recommendations for laboratory analysis in the diagnosis and management of diabetes mellitus" Diabetes care vol. 34,6 (2011): e61-99.

HOW TO USE IT

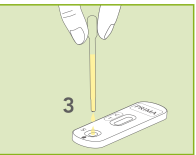
1) The sample is the first urine of the day collected in a clean cup, not provided with the kit.



2) Press the pipette bulb and dip the pipette tip into the collected urine. Release the bulb to collect the urine sample.



3) Deposit 3 drops of sample into the well (S) indicated on the cassette and wait 5 minutes before reading the result.



CONTENT: 1 sealed aluminium pouch containing: 1 test cassette and 1 desiccant bag; 1 plastic pipette for sampling and 1 instructions for use leaflet.

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(\land)	VIDEO	REF	DESCRIPTION	NUMBER OF TESTS	SHELF LIFE
C	TUTORIAL	820027-1	KIDNEYS ALBUMIN TEST	1 TEST	30 MONTHS





IRON FER TEST

Self-test for the detection of ferritin levels in whole blood samples



IRON STORAGE & FERRITIN

Iron is an essential metal for our bodies and is crucial for transporting oxygen in the blood, for cell multiplication and to build the structure of tissues and organs. However, in excessively high levels it is toxic for the body. For this reason, every one of us has a system for taking up iron from the external environment (e.g. via a diet rich in iron-containing foods) and storing it in cells in a way that is not excessive (and therefore not toxic).

Ferritin is the protein responsible for this storage function. The level of ferritin is an excellent indicator of the amount of iron available to the body. Low levels of this protein in the blood are an indication of depleted iron stores, a condition that precedes the development of anaemia. A decrease can be caused by pregnancy, haemorrhages, alterations in iron uptake.

WHO ARE THE INTENDED USERS

Everyone over the puberty age (\approx 15 years). Before normal levels are higher than the decision value of the cut-off level of this test.

WHY - BENEFITS

Checking the normality of the Ferritin value is a useful tool for verifying the iron deficiency anaemia.

TEST PRINCIPLE

IRON FER TEST is an immunochromatographic assay which detects the protein Ferritin thanks to special monoclonal gold-conjugate antibodies embedded to test strip.

TECH SPECS

CUT-OFF	SENSITIVITY	SPECIFICITY	OVERALL ACCURACY
30 ng/mL	85,20%	100,00%	96,60%

Performance data obtained by clinical study with 120 participants enrolled. Biokit Quantex Ferritin has been utilized as reference method.

CLINICAL EVIDENCES

1. Wick M, Pingerra W, Lehmann P, Iron metabolism: diagnosis and therapy of anemias, 5th ed, Vienna, New York: Springer Verlag, 2003; p. 151.

2. Worwood M. The laboratory assessment of iron status – an update. Clin Chim Acta 1997; 259: 3-23.

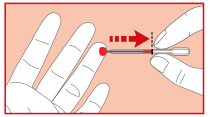
3. Kaltwasser JP, Werner E. Diagnosis and clinical evaluation of iron overload. Baillieres Clin Haematol 1989; 2; 363-89.

4. Baynes RD, Cook JD. Current issues in iron deficiency. Curr Opin Hematol 1996; 3:145-9.

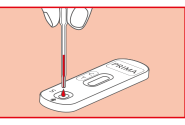
5. Lee MH, Means RT Jr. Extremely elevated serum ferritin levels in a university hospital: associated diseases and clinical significance. Am J Med 1996; 98: 566-71.

HOW TO USE IT

1) Take a blood sample after pricking the finger.



2) Deposit the sample into the specimen well of the cassette.



3) Add 2 drops into the well and wait 5 minutes before reading the result.



CONTENT: 1 sealed aluminium pouch containing: 1 test device and 1 desiccant bag; 1 transparent plastic bag containing a pipette for blood collecting; 1 vial with dropper containing the diluent; 2 sterile lancets for blood sampling; 1 alcohol swab and 1 instructions for use leaflet.

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	REF	DESCRIPTION	NUMBER OF TESTS	SHELF LIFE	
C	TUTORIAL	800025-1	IRON FER TEST	1 TEST	30 MONTHS





ALLERGY IgE TEST

Self-test for the determination of total class E Immunoglobulins (IgE) in whole blood samples



ALLERGIES AND IgE

Immunoglobulins E (IgE) are the class of antibodies that are most involved in allergic reactions. In normal conditions, IgE are present in blood in a really low quantity.

However, when they come in contact with the allergens, their concentration increases and they can trigger the cascade of reactions leading to the appearance of the main symptoms of an allergic reaction (such as skin erythema, itchiness, sneezing repeatedly, and, in some cases, anaphylactic shock).

This is why it is important to check if IgE levels are above normal, especially in people with a family history of allergies.

WHO ARE THE INTENDED USERS

ALLERGY IGE TEST is recommended to everyone with allergic reaction symptoms, such as itching and/or reddened skin, watery eyes and sneezing repeatedly.

WHY - BENEFITS

Checking the presence of IgE antibodies is useful to verify a potential general allergy reaction.

TEST PRINCIPLE

ALLERGY IGE TEST is an immunochromatographic test that detects the presence of IgE antibodies in the blood in concentrations above average, using special monoclonal antibodies conjugated with gold and integrated into the reactive strip.

TECH SPECS

CUT-OFF	SENSITIVITY	SPECIFICITY	OVERALL ACCURACY
100 IU/mL	93,42%	91,42%	92,46%

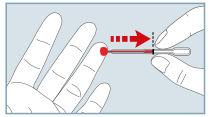
Performance data obtained by clinical study with 146 participants enrolled. Immunocap Total IgE Thermofisher has been utilized as reference method.

CLINICAL EVIDENCES

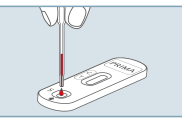
 Martins T.B., Bandhauer M.E., Bunker A. M., Roberts W. L.: "New childhood and adult reference intervals for total IgE." Journal of Allergy and Clinical Immunology. 2014;133(2):589-91.
 Chang M.L., Cui C., Liu Y.H., Pei L.C., Shao B.: "Analysis of total immunoglobulin E and specific immunoglobulin E of 3,721 patients with allergic disease". Biomed Rep. 2015 Jul;3(4):573-577.
 I. Leonard Bernstein, MD; James T. Li, et Al.: "Annals of allergy, asthma, & immunology." March 2008; Vol 100, num 3, supplement 3.

HOW TO USE IT

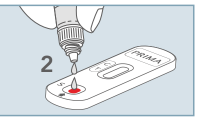
1) Take a blood sample after pricking the finger.



2) Deposit the sample into the specimen well of the cassette.



3) Add 2 drops into the well and wait 5 minutes before reading the result.



CONTENT: 1 sealed aluminium pouch containing: 1 test device and 1 desiccant bag; 1 transparent plastic bag containing a pipette for blood collecting; 1 vial with dropper containing the diluent; 2 sterile lancets for blood sampling; 1 alcohol swab and 1 instructions for use leaflet.

\bigwedge	VIDEO	REF	DESCRIPTION	NUMBER OF TESTS	SHELF LIFE
C	TUTORIAL	800018-1	ALLERGY IGE TEST	1 TEST	36 MONTHS







VITAMIN D TEST

Rapid self-test for the semi-quantitative detection of Vitamin D in human whole blood



DEFICIENT 0-10 (ng/mL) 0-25 (nmol/L)

- INSUFFICIENT 10-30 (ng/mL) 25-75 (nmol/L)
- SUFFICIENT 30-100 (ng/mL) 75-250 (nmol/L)

EXCESS

>100 (ng/mL) >250 (nmol/L)

VITAMIN D

Vitamin D refers to a group of fat-soluble secosteroids responsible for increasing intestinal absorption of calcium, iron, magnesium, phosphate and zinc. Vitamin Disproduced by skin exposed to ultraviolet B radiation or obtained from dietary sources, including supplements. Average daily vitamin D intake in the population and current dietary reference intake values are often inadequate to maintain optimal vitamin D levels. Virtually every cell in our body has receptors for Vitamin D, meaning that they all require "sufficient" level of Vitamin D for adequate functioning. Vitamin deficiency has been linked to various serious diseases: osteoporosis, multiple sclerosis, cardiovascular diseases, pregnancy complications, diabetes, strokes, autoimmune diseases, infectious diseases, etc.

WHO ARE THE INTENDED USERS

VITAMIN D TEST is intended for persons commonly at risk for vitamin D deficiency, such as people who get: tired and fatigued easily, inadequate sun exposure, limited oral intake, or have impaired intestinal absorption.

WHY - BENEFITS

VITAMIN D TEST helps understanding whether Vitamin D levels are sufficient, insufficient or deficient, allowing the person to take precautions in time.

TEST PRINCIPLE

VITAMIN D TEST is an immunochromatographic assay that, thanks to specific antibodies, is able to detect the levels of 25-hydroxyvitamin D in blood sample and to highlight its concentration thanks to the intensity of the test line (T) obtained.

TECH SPECS

CUT-OFF	RANGE OF MEASUREMENTS	OVERALL ACCURACY
30 ± 4 ng/mL	10-100 ng/mL	94,40%

Performance data obtained by comparing the test with another EC marked Vitamin D Rapid Test.

CLINICAL EVIDENCES

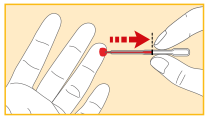
1. Grant WB, Holick MF (June 2005). Benefits and requirements of vitamin D for optimal health: a réview. Alternative Medicine Review.10 (2): 94–111.

2. Moyad MA. Vitamin D: a rapid review. DermatolNurs. 2009, 21:25-30

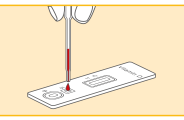
3. Holick MF (March 2006). "High prevalence of vitamin D inade-quacy and implications for health". Mayo Clinic Proceedings. 81 (3): 353–73.

HOW TO USE IT

1) Take a blood sample after pricking the finger.



2) Deposit the sample into the specimen well of the cassette.



3) Add 2 drops into the well and wait 10 minutes before reading the result.



CONTENT: 1 sealed aluminum pouch containing: 1 test device and 1 dessicant bag; 1 transparent plastic bag containing a pipette for blood collection; 1 vial containing the diluent; 1 color card; 2 sterile lancets for blood sampling; 1 alcohol swab and 1 instructions for use leaflet.

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(\land)	VIDEO	REF	DESCRIPTION	NUMBER OF TESTS	SHELF LIFE
C	TUTORIAL	100066-1	VITAMIN D TEST	1 TEST	24 MONTHS



00066-1_LEAFLET_02_1.8 05/2024





DRUG TEST

Self-test for the detection of 6 drugs in urine: Morphine, Amphetamine, Cocaine, Marijuana, Methamphetamine and Methadone



DETECTED SUBSTANCES

TEST	Cut-off (ng/mL)	Time permanence in urine after exposure
Amphetamine (AMP1000)	1000	1-3 days for a single administration, up to 9 days in chronic users
Cocaine (COC 300)	300	1-3 days for a single administration, up to 3 weeks in chronic users
Marijuana	50	single administration (smoking): 30 h on average, up to 4 days single administration (oral intake): up to 6 days occasional use (once or twice a week): up to 30 days regular consumption: up to 3 months
Methadone (MTD 300)	300	1,5 to 3 days
Methamphetamine (MET 500)	500	1 to 4 days for a single administration, up to 5 days in chronic users
Morphine (MOP 300)	300	10-55 h for a single administration, up to 11 days in chronic users

WHO ARE THE INTENDED USERS

DRUG TEST can be used by everyone who wants to check the presence of the above mentioned substances in urine.

TEST PRINCIPLE

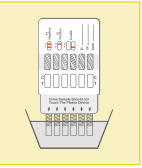
DRUG TEST is an immunochromatographic assay that can be performed without the use of an instrument. The test utilizes monoclonal antibodies to selectively detect levels of specific drugs in urine.

TECH SPECS

	CUT-OFF	SENSITIVITY	SPECIFICITY	OVERALL ACCURACY
AMP	1000 ng/mL	99,90%	99,90%	99,90%
MET	500 ng/mL	99,90%	99,90%	99,90%
COC	300 ng/mL	99,90%	99,90%	99,90%
MOP	300 ng/mL	99,90%	99,90%	99,90%
MTD	300 ng/mL	99,90%	99,90%	99,90%
THC	50 ng/mL	99,09%	99,90%	99,90%

HOW TO USE IT

1) Open the cap, with the arrow pointing toward the urine specimen, dip the test panel vertically in the urine specimen for at least 10 to 15 seconds. 2) Replace the cap and leave the test panel on a non-absorbent flat surface. 3) Wait 5 minutes before reading the result.



CONTENT: 1 sealed aluminium pouch containing: 1 test panel and 1 desiccant bag; 1 instructions for use leaflet.



VIDEO	REF	DESCRIPTION	NUMBER OF TESTS SHELF LIFE	
TUTORIAL	19806-1	DRUG TEST	1 TEST	24 MONTHS



EARLY PREGNANCY TEST

Rapid self-test for the early detection of the hCG in human urine samples (10 mIU/mL)



PREGNANCY AND hCG

In the very early stages of pregnancy, the fertilized egg is transported from the tubes to the uterus where, around seven days later, it lurks. Precisely at this stage, the tissue that will form the placenta begins to produce the human Chorionic Gonadotropin, also called hCG hormone, essential for the correct development of pregnancy.

The quick appearance and rapid increase in the concentration of the hCG hormone in the urine make it an excellent indicator for the early confirmation of pregnancy.

WHO ARE THE INTENDED USERS

Women who want to identify early stages of pregnancy.

WHY - BENEFITS

The test detects the hCG hormone in urine at a concentration of 10 mIU/mL, therefore the test can be performed 3 days before the period is due. So in this way it is possible to early identify a pregnancy.

TEST PRINCIPLE

EARLY PREGNANCY TEST is an immunochromatographic assay for the early detection of hCG hormone in urine by using a combination of one monoclonal-dye conjugate and another one monoclonal-solid phase antibody to selectively identify hCG in test samples.

TECH SPECS

CUT-OFF	SENSITIVITY	SPECIFICITY	OVERALL ACCURACY
10 mIU/mL	100%	100%	100%

Performance data obtained by clinical study with 47 participant enrolled. Clearblue Test Early Pregnancy has been used as reference method.

CLINICAL EVIDENCES

1. Batzer FR. "Hormonal evaluation of early pregnancy", Fertil. Steril. 1980; 34(1): 1-13 2.

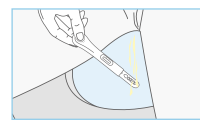
2. Snjder J.A., Haymonth F., Parvin C.A. et al. "Diagnostic considerations in the measurement of Human

Chorionic Gonadotrophin in aging women" Clin Chem 2005; 51: 1830 – 1835

3. Gnoth C., Johnson S.: "Strips of hope: accuracy of home pregnancy tests and new development"

HOW TO USE IT

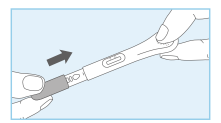
1) Remove the protective cap and place the absorbing tip right under the flow of urine for at least 10 seconds.



2) If easier, collect the urine in a clean, dry, residue-free container and dip the absorbing tip into the urine for 10 seconds.



3) Place the protective cap back onto the absorbing tip.Wait 5 minutes before reading the result.



CONTENT: 1 or 2 sealed aluminium pouches containing each: 1 spoon device and 1 desiccant bag; 1 instructions for use leaflet.

REF	DESCRIPTION	NUMBER OF TESTS	SHELF LIFE
820010-1	EARLY PREGNANCY TEST	1 TEST	
820010-2	EARLY PREGNANCY TEST	2 TESTS	30 MONTHS



OVULATION LH TEST

Self-test for the detection of luteinizing hormone (LH) in urine



LH AND OVULATION

FERTILITY AREA FEMALE FERTILITY

During the entire childbearing age, the period of time between puberty and menopause, women reproductive system meets monthly a series of structural and functional important changes. This happens because of the continuous changes in estrogen levels and progesterone, hormones that are regulated by the Hypothalamus-Hypophysis-Ovary axis. Their concentrations are different during the 4 phases of menstrual cycle: menstruation, follicular phase, ovulation and luteal phase. During the ovulation phase, which occurs around the 12th/13th day of the menstrual cycle, the level of estrogen in the blood is so high that the hypophysis starts the production of the Luteinizing Hormone (LH), thus determining the breakdown of the follicle and the release of the egg cell, ready to be fertilized. The egg cell remains fertilizable up to the next 24 hours.

WHO ARE THE INTENDED USERS

Women who want to conceive.

WHY - BENEFITS

Measuring the levels of hormone LH in the urine is an effective method to optimize the possibility of conception.

TEST PRINCIPLE

OVULATION LH TEST is an immunochromatographic assay that detects LH through special goldconjugated monoclonal antibodies included in the reactive strip.

TECH SPECS

CUT-OFF	SENSITIVITY	SPECIFICITY	OVERALL ACCURACY
25 mIU/mL	100,00%	97,14%	98,11%

Performance data obtained by clinical study with 53 participants enrolled. iXensor Ovulation LH test (Immunochromatographic Lateral Flow) has been utilized as reference method.

CLINICAL EVIDENCES

1. Su HW, Yi YC, Wei TY, Chang TC, Cheng CM "Detection of ovu-

1. Survey, M. C., Wein, Chang M. C., Cheng C.W. Detection Over-lation, a review of currently available method" Bioeng Transl Med. 2017 May 16;2(3):238-246.
2. Leiva RA, Bouchard TP. Abdullah SH, Ecochard R.: "Urinary Luteinizing Hormone Tests: Which concentration threshold best predict ovulation?" Front public health 2017, 5: 320.

HOW TO USE IT

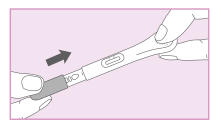
1) Remove the protective cap and place the absorbing tip right under the flow of urine for at least 10 seconds.



2) If easier, collect the urine in a clean, dry, residue-free container and dip the absorbing tip into the urine for 10 seconds.



3) Place the protective cap back onto the absorbing tip.Wait 5 minutes before reading the result.



CONTENT: 5 sealed aluminium pouches containing each: 1 spoon device and 1 desiccant bag; 1 instructions for use leaflet.

\bigcap	VIDEO	REF	DESCRIPTION	NUMBER OF TESTS	SHELF LIFE
E	TUTORIAL	820050-5	OVULATION LH TEST	5 TESTS	30 MONTHS





MENOPAUSE FSH TEST

Self-test for the determination of the Follicle Stimulating Hormone - FSH in urine



MENOPAUSE AND FSH

Menopause is defined as the time when there has been no menstrual periods for 12 consecutive months and no other biological or physiological cause can be identified. The menopause is caused by a modification of the hormonal balance in woman's body. This change arises generally when close to 45 years old. But true signs are observed around 55 years old. An irregular duration of periods is an early indication of the menopause beginning (peri-menopause). Menopause is the consequence of the end of ovary follicle life-cycle leading to increase of Follicle Stimulating Hormone (FSH) circulating level. Concentration of FSH increases during menopause passing from values less than 20 mUI/mL up to approximately 80 mUI/mL.

WHO ARE THE INTENDED USERS

WOMEN > 45 years.

WHY - BENEFITS

Checking the presence of high levels of FSH is a useful tool for verifying a potential ongoing menopause status.

TEST PRINCIPLE

MENOPAUSE FSH TEST is an immunochromatographic test that detects FSH through special goldconjugated monoclonal antibodies included in the reactive strip.

TECH SPECS

CUT-OFF	SENSITIVITY	SPECIFICITY	OVERALL ACCURACY	
25 mIU/mL	100,00%	86,90%	94,00%	

Performance data obtained by clinical study with 100 participants enrolled. Roche Cobas 8000 has been utilized as reference method.

CLINICAL EVIDENCES

1. Robert J. Norman "Fertility testing" Australian Prescriber, 2002;25:38-401

2. Buckler H. "The menopause transition: endocrine changes and clinical symptoms." J Br Menopause Soc. 2005 Jun;11(2):61-5.

3. National Collaborating Centre for Women's and Children's Health (UK) "Menopause: Full Guideline". 2015 Nov. National Institute for Health and Care Excellence: Clinical Guidelines.

HOW TO USE IT

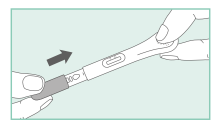
1) Remove the protective cap and place the absorbing tip right under the flow of urine for at least 10 seconds.



2) If easier, collect the urine in a clean, dry, residue-free container and dip the absorbing tip into the urine for 10 seconds.



3) Place the protective cap back onto the absorbing tip.Wait 5 minutes before reading the result.



CONTENT: 2 sealed aluminium pouches containing each: 1 spoon device and 1 desiccant bag; 1 instructions for use leaflet.

\bigcap	VIDEO	REF	DESCRIPTION	NUMBER OF TESTS	SHELF LIFE
	820011-2	MENOPAUSE FSH TEST	2 TESTS	30 MONTHS	



SPERM TEST

Rapid self-test for in-vitro qualitative estimation of sperm concentration in human semen



MALE INFERTILITY

FERTILITY AREA MALE FERTILITY

Male infertility can be an health issue for men and is primarily responsible for inability to conceive after 1 year of regular, unprotected intercourses. Particularly, male factor infertility affects nearly 50% of infertile worldwide couples who want to conceive and to require empirical therapy. Sperm count is considered lower than normal if you have fewer than 15 million sperm per milliliter of semen.

WHAT IS SPERM TEST

SPERM TEST is designed to be used for in vitro qualitative estimation of the sperm concentration of human semen. The test will determine if the number of sperms is adequate for conception, subject to female partner's ovulation in time. A low sperm concentration would indicate less likelihood of conception.

WHO ARE THE INTENDED USERS

Men who want to check their sperm concentration.

WHY - BENEFITS

Checking the sperm concentration is a useful tool for verifying the male fertility status.



SPERM TEST is biochemical assay for in vitro qualitative estimation of sperm concentration in human semen. Sperm cells are trapped on the first surface layer of the membrane, while a staining solution is used to dye sperm cells. The darker is the colour of well A, the higher is the sperm concentration. If the colour of well A is lighter than the standard colour of reference well B, it means that the concentration of sperm is less than 15 million/mL. If the colour of well A test is darker than the standard colour of reference well B, it means that the sperm concentration is greater than 15 million/mL.

TECH SPECS

CUT-OFF	SENSITIVITY	SPECIFICITY	OVERALL ACCURACY
15 million/mL	98,10%	99,50%	99,10%

Performance data obtained by clinical study with 236 participants enrolled. Sperm Concentration Rapid Test (Colorimetric) has been utilized as reference method.

CLINICAL EVIDENCES

1. Centola G.M.: Semen assessment. Urol Clin North Am. 2014 Feb;41(1):163-7. doi: 10.1016/j.ucl.2013.08.007. Epub 2013 Sep 13. 2. Cooper T.G., Noonan E., von Eckardstein S., Auger J., Baker H.W., Behre H.M., Haugen T.B., Kruger T., Wang C., Mbizvo M.T., Vogelsong K.M. World Health Organization reference values for human semen characteristics. Hum Reprod Update. 2010 May-Jun;16(3):231-45.

HOW TO USE IT

1) Collect the semen directly into the collection cup. Shake it evenly in the collection cup and leave it to stand for 15 minutes until the semen liquefies.







3) Add three drops of the blue staining solution to well A. Let it soak for 1-2 minutes.

4) Apply two drops of the transparent washing solution to well A, and let it soak for 1-2 minutes, then read the results immediately by comparing the colour of A to B.



CONTENT: 1 sealed aluminium pouch containing: 1 test device, 1 desiccant bag and 1 pipette; 1 vial with dropper tip containing the staining solution; 1 vial with dropper tip containing the washing solution; 1 collection cup and 1 instructions for use leaflet.



ΕO	REF	DESCRIPTION	NUMBER OF TESTS	SHELF LIFE
FORIAL	100082-1	SPERM TEST	1 TEST	24 MONTHS





0483

WHAT IS AMNIOCHECK

AMNIOCHECK is designed to detect any minute (100 microlitre) leak of amniotic fluid, differentiating it from urine by detecting the pH level.

WHO ARE THE INTENDED USERS

- Gynecologists' clinics or Delivery rooms: for amniotic fluid leakage screening or verification, especially for non-continuous leakage.
- Pregnant women at home:

women with normal pregnancies experiencing unexplained wetness as well as by those at high risk of Premature Rupture of Membrane (PROM) and Preterm Premature Rupture of Membrane (PPROM).

Availabe for home use and prefessional use.

WHY - BENEFITS

Non invasive method, easy to use; early diagnosis of PROM and PPROM; reduction of unnecessary speculum examination; reduction of time spent in hospital; reduced risk of infection from speculum examination (particularly if repeat examinations are required); incidental detection of possible vaginal infection; reduction in staff time and hospital bed use.

AMNIOCHECK

Amniotic leak self-test detector



TEST PRINCIPLE

AMNIOCHECK changes colours when it comes in contact with fluid of pH levels greater than or equal to 6,5 units. Amniotic fluid pH levels are greater than 6,5.

The panty liner is specific to vaginal fluid, and if contaminated with urine, in the range of pH 5,5-7,0, will not give a positive result (combination of the chemical formulation and panty liner composition reduces interference from urine and therefore increase specificity).

TECH SPECS

CUT-OFF	SENSITIVITY	SPECIFICITY	OVERALL ACCURACY
pH > 6,5	97,10%	96,90%	PPV 96,10% NPV 97,67%

Performance data obtained by clinical studies conducted in 2016-2017 (5 American centres and 1 Israelian centre) with 232 pregnant women enrolled. AMNIOCHECK has been compared to the standard clinical diagnosis protocol.

CLINICAL EVIDENCES

The National Institute for Health and Care Excellence (NICE, UK) has conducted evaluated this panty liner as "sufficiently accurate to exclude amniotic fluid leak as a cause of wetness in pregnancy, [...] the use of the panty liner could generate costs savings and avoid the discomfort of unnecessary speculum examinations, [...] its use should be encouraged."

Bibliography: 1) Bornstein J, Geva A, Salt I, Fait V, Schoenfeld A, Shoham HK, Sobel J. Nonintrusive diagnosis of premature ruptured amniotic membranes using a novel polymer. Am J Perinatol. 2006 Aug; 23(6):351-4. 2) Bornstein J, Ohel G, Sorokin Y, Reape KZ, Shnaider O, Kessary-Shoham H, Ophir E. Effectiveness of a novel home-based testing device for the detection of rupture of membranes. Am J Perinatol. 2009 Jan;26(1):45-50.

HOW TO USE IT

Attach the panty liner to the underwear; the panty liner has a central polymer-embedded strip that turns blue-green on contact with fluid. Wear the panty liner as a normal panty liner till feel wetness (do not exceed 12 hours). At the first sense of vaginal wetness, remove the panty liner and wait up to 15 minutes and check for a colour change on the panty liner.

The detection panty liner is specific to vaginal fluid, and if contaminated with urine it will not give a positive result.



CONTENT: 5 aluminum pouches containing 1 panty liner each; 1 instructions for use leaflet.

	VIDEO	REF	DESCRIPTION	NUMBER OF TESTS SHELF LIFE	
	/ TUTORIAL	100088-5	AMNIOCHECK	5 TESTS	48 MONTHS



GIARDIA PET TEST

Rapid test for the qualitative identification of Giardia lamblia antigens in animal faeces



GIARDIA

Giardia lambia is a microscopic protozoan contracted by oral ingestion. It attacks the inner wall of animal's small intestine and interferes with the normal fats and carbohydrates absorption during digestion.

WHO ARE THE INTENDED USERS

GIARDIA PET TEST is recommended to everyone who wants to check their pet's health status. More common symptoms are: nausea, diarrhea, fever and weight loss.

WHY - BENEFITS

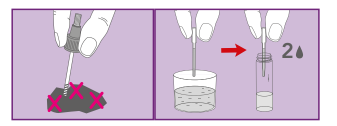
It is important to diagnose Giardiasis as soon as possible. The disease, if not treated, can lead to chronic debilitation with anorexia, progressive culling, slimming and growth retardation.

TEST PRINCIPLE

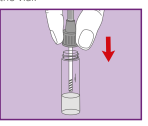
Giardia Test is a lateral flow immunochromatographic assay which detects the Giardia antigen using highly specific monoclonal antibodies.

HOW TO USE IT

1) Collect the sample and dip the stick into feces in different points.



2) Put the stick in the stool extraction liquid and shake the vial.



3) Apply 2 drops of the diluent stool sample. Wait 10 minutes and read the result.



CONTENT: 1 sealed aluminum pouch containing: 1 GIARDIA PET TEST and 1 desiccant bag; 1 transparent plastic bag containing a pipette for specimen collection; 1 vial with green screw cap with collection stick and dripper; instructions for use.

REF	DESCRIPTION	NUMBER OF TESTS	SHELF LIFE
PET100-1	GIARDIA PET TEST	1 TEST	24 MONTHS



VITAMIN D **RAPID TEST**

Rapid test for the semi-quantitative detection of 25-hydroxyvitamin D in human whole blood



TECH SPECS



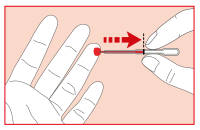
PRINCIPLE OF THE TEST

VITAMIN D RAPID TEST is a rapid immunochromatographic assay that is able to detect the levels of 25-hydroxyvitamin D in blood sample and highlight its concentration thanks to the different intensity of the Test Line (T) obtained.

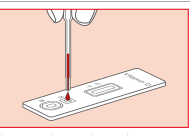
WHO ARE THE INTENDED USERS

The device is intended to be used by healthcare professionals as a preliminary diagnostic assay. The result obtained with the test indicates whether Vitamin D levels in whole blood are sufficient, insufficient, deficient or excessive allowing to take precautions in time.

HOW DOES THE TEST WORK?



1. Take a blood sample after pricking the finger, using the provided pipette.



2. Deposit the sample into the specimen well (S) of the cassette.



3. Add 2 drops of diluent in the buffer well (B) and wait 10 minutes before reading the result.

BIBLIOGRAPHY

- 1. Holick MF (March 2006). "High prevalence of vitamin D inadequacy and implications for health." Mayo Clinic Proceedings. 81 (3): 353–73. 2. Eriksen EF, Glerup H (2002). Vitamin D deficiency and aging: implications for general health and osteoporosis. Biogerontology. 3 (1-2): 73-7. 3. Grant WB, Holick MF (June 2005). Benefits and requirements of vitamin D for optimal health: a review. Alternative Medicine Review.10 (2): 94-111.
- 4. Moyad MA. Vitamin D: a rapid review. DermatolNurs. 2009, 21:25-30

REF	DESCRIPTION	NUMBER OF TESTS	SHELF LIFE	
100066IP-20P	VITAMIN D RAPID TEST	20 TESTS	24 MONTHS	



STREP A RAPID TEST

Rapid test for the detection of group A B-hemolytic Streptococcus in pharyngeal swab



TECH SPECS



PRINCIPLE OF THE TEST

STREP A RAPID TEST is a rapid immunochromatographic assay that detects the presence of specific Streptococcus A antigens in throat swab samples by using special monoclonal antibodies.

WHO ARE THE INTENDED USERS

The device is intended to be used by healthcare professionals in order to quickly determine whether a sore throat is caused by a Strep A or other germs (usually viruses), which don't require antibiotic treatments. Useful especially with children.



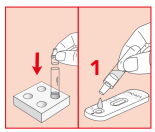
HOW DOES THE TEST WORK?

1. Collect a throat sample using the provided material.

2. Insert the absorbent tip into the extraction tube, previously placed on the workstation. Add R1 and R2 solutions.



3. Mix the solutions and leave the swab inside the tube for 2-5 minutes. Then remove it and squeeze the swab tip to collect as much liquid as possible.



4. Apply 1 drop of the extracted specimen and wait 5 minutes before reading the result.

BIBLIOGRAPHY

- Banerjee S, Ford C. "Rapid Tests for the Diagnosis of Group A Streptococcal Infection: A Review of Diagnostic
 Test Accuracy, Clinical Utility, Safety, and Cost-Effectiveness ". Ottawa (ON): Canadian Agency for Drugs and Technologies in Health; 2018 May, CADTH Rapid Response Reports.
 Centers for Disease Control and Prevention. Adult appropriate antibiotic use summary. http://www.cdc.gov/getsmart/community/materials-references/print-materials/hcp/
 adult-approp-summary.pdf. Accessed March 8, 2016.
 Lor C, Bjerrum L, Munck A., Cots JM., Henrández S., Moragas A.; HAPPY AUDIT Investigators. "Access to point-of-care tests reduces the prescription of antibiotics
 among antibiotic-requesting subjects with respiratory tract infections." Respir Care. 2014 Dec.

REF	DESCRIPTION	NUMBER OF TESTS	SHELF LIFE
800060IP-20P	STREP A RAPID TEST	20 TESTS	30 MONTHS



KI SCREENING TEST Canine NGAL Detection

The first and only rapid test for semi-quantitative detection of NGAL in urine for early screening of kidney damage in dogs



ANALYTICAL PERFORMANCES



PRINCIPLE OF THE TEST

KI SCREENING TEST is a lateral flow immunochromatographic device, for veterinary use, for semi-quantitative detection of NGAL in dog urine samples.

WHO ARE THE INTENDED USERS

The device is intended to be used by healthcare professionals in order to quickly determine the concentration of NGAL in dog urine, which is directly proportional to the severity of kidney damage and indicative of potential clinical settings.



HOW DOES THE TEST WORK?

1. Using the pipette provided, collect the urine sample.

BIBLIOGRAPHY



2. Deposit 2 drops of sample into the diluent bottle.



3. Close the diluent bottle with the cap and mix by gently turning the bottle upside down.



4. Add 3 drops of the mixed solution to the test cassette and wait 10 minutes before reading the result.

PRIMA

 F. Tagliasacchi, Abstract: Urinary neutrophil gelatinase-associated lipocalin (NGAL): A rapid lateral flow test in canine practice. Oral communication at the 33^{ed} ECVIM-CA Congress, 2023, Barcellona. Davis J, Raisis AL, Miller DW, Hosgood GL, Rossi G. Analytical validation and reference intervals for a commercial multiplex assay to measure five novel biomarkers for acute kidney injury in canine urine. Res Vet Sci. 2021 Oct; 139:78-86. Lee YJ, Hu YY, Lin YS, Chang CT, Lin FY, Wong ML, Kuo-Hsuan H, Hsu WL. Urine neutrophil gelatinase-associated lipocalin (NGAL) as a biomarker for acute canine kidney 		injury, BMC Vet Res. 2012 Dec 28;8:248. 4. Nabity M, Hokamp J. Urinary Biomarkers of Kidney Disease in Dogs and Cats. Vet Clin North Am Small Anim Pract. 2023 Jan;53(1):53-71. doi: 10.1016/j.cvsm.2022.07.006. Epub 2022 Oct 19. PMID: 36270837. 5. Steinbach S, Weis J, Schweighauser A, Francey T, Neiger R. Plasma and urine neutrophil gelatinase-associated lipocalin (NGAL) in dogs with acute kidney injury or chronic kidney disease. J Vet Intern Med. 2014 Mar-Apr;28(2):264-9. doi: 10.1111/jvim.12282.			022.07.006. Epub Ind urine neutrophil or chronic kidney	
	REF	DESCRIPTION		NUMBER OF TESTS	SHELF LIFE	
	800087-5	KI SCREENING TEST		5 TESTS	18 MONTHS	primalabsa.ch









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