Urinalysis Reagent Strips Package Insert (Urine)

For rapid detection of multiple analytes in human urine For in vitro diagnostic use only

of one or more of the following analytes in urine: Ascorbic acid, Glucose, Bilirubin, and Leukocytes. The Urinalysis Reagent Strips (Urine) are for single use Ketone (Acetoacetic acid), Specific Gravity, Blood, pH, Protein, Urobilinogen, Nitrite reagent areas are affixed. The test is for the qualitative and semi-quantitative detection The Urinalysis Reagent Strips (Urine) are firm plastic strips onto which several separate

analyte(s) and color blocks on the color chart for results Refer to kit box label for the specific analyte(s) listed, and compare to the appropriate professional near-patient (point-of-care) and centralized laboratory locations.

in the diagnosis and monitoring of metabolic or systemic diseases that affect kidney function, endocrine disorders and diseases or disorders of the urinary tract. 12 Urinalysis Reagent Strips (Urine) can be used in general evaluation of health, and aids an indicator of health or disease, and as such, is a part of routine health screening. The blood composition is altered to a significant extent. Urinalysis is a useful procedure as Urine undergoes many changes during states of disease or body dysfunction before (SUMMARY)

[PRINCIPLE AND EXPECTED VALUES]

Patients with adequate diet may excrete 2-10 mg/dL daily. After ingesting large Ascorbic acid: This test involves decolorization of Tillmann's reagent. The presence of ascorbic acid causes the color of the test field to change from blue-green to orange. amounts of ascorbic acid, levels can be around 200 mg/dL

and hydrogen peroxide in the presence of glucose oxidase. The hydrogen peroxide oxidase, peroxidase and chromogen. Glucose is first oxidized to produce gluconic acid Glucose: This test is based on the enzymatic reaction that occurs between glucose glucose may be excreted by the kidney.3 Glucose concentrations as low as 100 mg/Dl green to brown. Glucose should not be detected in normal urine. Small amounts of which the chromogen is oxidized determines the color which is produced, ranging from reacts with potassium iodide chromogen in the presence of peroxidase. The extent to may be considered abnormal if results are consistent

dichloroaniline in a strongly acidic medium. Varying bilirubin levels will produce a pinkish-tan color proportional to its concentration in urine. In normal urine, no bilirubin is present in the urine specimen, and are possibly masking the bilirubin reaction. blocks shown on the color chart) may indicate that bilirubin-derived bile pigments are further investigation. Atypical results (colors different from the negative or positive color detectable by even the most sensitive methods. Even trace amounts of bilirubin require Bilirubin: This test is based on azo-coupling reaction of bilirubin with diazotized

concentration before serum ketones are elevated. carbohydrate metabolism situations, ketones appear in the urine in excessively high ketone levels may occur in urine during physiological stress conditions such as fasting, pregnancy and frequent strenuous exercise. *4 In starvation diets, or in other abnormal purple color for positive results. Ketones are normally not present in urine. Detectable to produce a color change ranging from light pink for negative results to a darker pink or Ketone: This test is based on ketones reacting with nitroprusside and acetoacetic acid

Specific Gravity: This test is based on the apparent pKa change of certain pretreated range from deep blue-green in urine of low ionic concentration polyelectrolytes in relation to ionic concentration. In the presence of an indicator, colors severe renal damage, the specific gravity is fixed at 1.010, the value of the glomerular with normal diets and fluid intake will have a specific gravity of 1.016-1.022.8 in cases of vary in specific gravity from 1.003-1.035.8 Twenty-four hour urine from healthy adults yellow-green in urine of increasing ionic concentration. Randomly collected urine may to green and

patients and clinical judgment is required in these specimens. urine of menstruating females. The significance of a trace reading varies among specimen should be examined further. Blood is often, but not invariably, found in the color development on the reagent area within 60 seconds is significant and the urine the reaction of diisopropylbenzene dihydroperoxide and 3,3,5,5'-tetramethylbenzidine Blood: This test is based on the peroxidase-like activity of hemoglobin which catalyzes The resulting color ranges from orange to green to dark blue. Any green spots or green

expected range for other normal urine specimens is pH 4.5-8, with an average result of covering the entire urinary pH range. Colors range from orange to yellow and green to blue. The expected range for normal urine specimens from newborns is pH 5.7.9 The pH: This test is based on a double indicator system which gives a broad range of colors

proteins (anions) as the indicator releases hydrogen ions to the protein. At a constant indicators where an indicator that is highly buffered will change color in the presence of Protein: This reaction is based on the phenomenon known as the "protein error" of pH

> from yellow to yellow-green for negative results and green to green-blue for positive results. 1-14 mg/dL of protein may be excreted by a normal kidney. 6 A color matching required to evaluate the significance of trace results any block greater than trace indicates significant proteinuria. Clinical judgment is pH, the development of any green color is due to the presence of protein. Colors range

0.2-1.0 mg/dL (3.5-17 µmol/L).8 A result of 2.0 mg/dL (35 µmol/L) may be of clinical and is a normal substance in urine. The expected range for normal urine with this test is pink color. Urobilinogen is one of the major compounds produced in heme synthesis p-diethylaminobenzaldehyde and urobilinogen in strongly acidic medium to produce a Urobilinogen: This test is based on a modified Ehrlich reaction between

negative bacteria in the urine. In an acidic medium, nitrite in the urine reacts with significance and the patient specimen should be further evaluated. collection. Retrieval of positive cases with the nitrite test ranges from as low as 40% in depending on how long the urine specimens were retained in the bladder prior to detectable in normal urine.9 The nitrite area will be positive in some cases of infection, couples with 1 N-(1-naphthyl) ethylenediamine to produce a pink color. Nitrite is not p-arsanilic acid to form a diazonium compound. The diazonium compound in turn Nitrite: This test depends upon the conversion of nitrate to nitrite by the action of Gram cases where little bladder incubation occurred, to as high as approximately 80% in cases where bladder incubation took place for at least 4 hours.

cleave a derivatized pyrazole amino acid ester to liberate derivatized hydroxyl pyrazole. using a fresh specimen from the same patient. Repeated trace and positive results are questionable clinical significance. When trace results occur, it is recommended to retest Normal urine specimens generally yield negative results. Trace results may be of This pyrazole then reacts with a diazonium salt to produce a beige-pink to purple color. Leukocytes: This test reveals the presence of granulocyte esterases. The esterases

[REAGENTS AND PERFORMANCE CHARACTERISTICS]

Based on the dry weight at the time of impregnation, the concentrations given may vary within manufacturing tolerances. The following table below indicates read times and

performance characteristics for each parameter Urobilinogen Ascorbic Reagent Leukocytes Gravity (KET) Bilirubin Glucose (GLU) (ASC Specific (BLO) (BIL) Acid (PRO) Protein (LEU) Nitrite (URO) E seconds Read seconds seconds seconds seconds Secolina seconds 30 seconds 30 30 60 45 40 60 60 60 120 60 sodium nitroprusside; buffer salt; buffer and non-reactive potassium 2,6-dichlorophenolindophenol; Detects ascorbic acid as low dihydroperoxide; buffer (TMB); ingredients; poly (methyl vinyl|1.030. Results correlate with glucose oxidase; peroxidase; tetrabromophenol blue; buffer bromthymol blue; non-reactive differentiation of pH ether/maleic 4-dichloroaniline diazoniun non-reactive ingredients ester; diazonium salt; buffer; 9-15 white blood cells Leu/µL and non-reactive ingredients non-reactive ingredients ethylenediamine; non-reactive non-reactive ingredients promthymol blue indicator, Determines urine derivatized pyrrole amino acid|Detects leukocytes as low as nethyl 3,3',5,5'-tetramethylbenzidine odium hydroxide p-arsanilic acid; N-(1-naphthyl -diethylaminobenzaldehyde; Composition and and red and diisopropylbenzene iodide; sodium non-reactive as 5-10 mg/dL (0.28-0.56 non-reactive gravity non-reactive anhydride); values obtained by refractive buffer salt; Permits and 50-100 mg/dL (2.5-5 mmol/L). Detects glucose as low as 5-10 low as 2.5-5 mg/dL (0.25-0.5 0.4-1.0 mg/dL (6.8-17 µmol/L) Detects bilirubin as low as low as 0.018-0.060 mg/dL or as 0.05-0.1 mg/dL in urine with Detects urobilinogen as low as 7.5-15 mg/dL (0.075-0.15 g/L) Detects free hemoglobin index method within ± 0.005. 0.2-1.0 mg/dL (3.5-17 µmol/L) Detects albumin as low content of < 50 mg/dL. specimens with ascorbic acid nmol/L) etects acetoacetic acid as in clinical urine a low specific gravity and less Detects sodium nitrite as low vithin the range of 5-9 han 30 mg/dL ascorbic acid. between 1.000 Description Ery/µL the 5 quantitative specific Agines un ne and

The performance characteristics of the Urinalysis Reagent Strips (Urine) have been

developed to be specific for the parameters to be measured with the exceptions of the are sensitivity, specificity, accuracy and precision. Generally, this test has been Interpretation of visual results is dependent on several factors: the variability of color interferences listed. Please refer to the Limitations section in this package insert

determined in both laboratory and clinical tests. Parameters of importance to the user

when the strip is read. Each color block on the chart corresponds to a range of analyte perception, the presence or absence of inhibitory factors, and the lighting conditions

[PRECAUTIONS

- · For in vitro diagnostic use only. Do not use after the expiration date
- The strip should remain in the closed canister until use.
- Do not touch the reagent areas of the strip.
- Discard any discolored strips that may have deteriorated.
- All specimens should be considered potentially hazardous and handled in the same
- manner as an infectious agent. The used strip should be discarded according to local regulations after testing.

STORAGE AND STABILITY

(2-30°C). Keep out of direct sunlight. The strip is stable through the expiration date printed on the canister label. Do not remove the desiccant. Remove only enough strips Store as packaged in the closed canister either at room temperature or refrigerated for immediate use. Replace cap immediately and tightly. DO NOT FREEZE. Do not use beyond the expiration date.

months. Stability may be reduced in high humidity conditions. Note: Once the canister has been opened, the remaining strips are stable for up to 3

[SPECIMEN COLLECTION AND PREPARATION]

possible. Do not centrifuge. The use of urine preservatives is not recommended. A urine specimen must be collected in a clean and dry container and tested as soon as testing cannot be done within an hour after voiding, refrigerate the specimen immediately and let it return to room temperature before testing

Prolonged storage of unpreserved urine at room temperature may result in microbial positive results with the protein test area. Urine containing glucose may decrease in pH proliferation with resultant changes in pH. A shift to alkaline pH may cause false as organisms metabolize the glucose.

affect protein (and to a lesser extent, specific gravity and bilirubin) test results. Contamination of the urine specimen with skin cleansers containing chlorhexidine may

Materials Provided Package insert

Materials Required But Not Provided

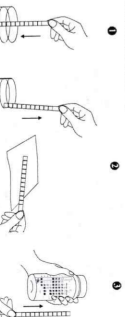
Timer

Specimen collection containers

Allow the strip, urine specimen, and/or controls to reach room temperature [DIRECTIONS FOR USE]

- (15-30°C) prior to testing .Remove the strip from the closed canister and use it as soon as possible Completely immerse the reagent areas of the strip in fresh, well-mixed urine and Immediately close the canister tightly after removing the required number of strip(s) immediately remove the strip to avoid dissolving the reagents. See illustration 1
- 2. While removing the strip from the urine, run the edge of the strip against the rim of to avoid mixing chemicals from adjacent reagent areas and/or soiling hands with urine. See illustration 2 below. bring the edge of the strip into contact with an absorbent material (e.g. a paper towel) the urine container to remove excess urine. Hold the strip in a horizontal position and
- 3. Compare the reagent areas to the corresponding color blocks on the canister label at the specified times. Hold the strip close to the color blocks and match carefully. See illustration 3 below.

be read using the Urine Analyzers. Refer to the Instruction Manual for details. Note: Results may be read up to 2 minutes after the specified times. Results may also



[INTERPRETATION OF RESULTS]

and repeat the test using a new strip. If the problem persists, discontinue using the strip printed on the canister label, compare results with known positive and negative controls immediately and contact your local distributor. are recommended: confirm that the strips have been tested within the expiration date nominal values. In the event of unexpected or questionable results, the following steps label. The color blocks represent nominal values; actual values will vary close to the Results are obtained by direct comparison of the color blocks printed on the canister

[QUALITY CONTROL]

for adequate standards of performance. positive and negative specimens/controls whenever a new test is performed, For best results, performance of reagent strips should be confirmed by testing known whenever a new canister is first opened. Each laboratory should establish its own goals or

[LIMITATIONS]

could be interpreted as false results. development on the test pad may be masked or a color reaction may be produced that abnormal urine color such as drugs containing azo dyes (e.g. Pyridium®, Azo Gantrisin®, Azo Gantrisin®, Furadantin®), and riboflavin.8 The color Note: The Urinalysis Reagent Strips (Urine) may be affected by substances that cause

Ascorbic acid: No interference is known.

mg/dL may cause false negative results for specimens containing a small amount of nalidixic acid). Sensitivity may be decreased in specimens with high specific gravity (> glucose (50-100 mg/dL). 1.025) and with ascorbic acid concentrations of ≥ 25 mg/dL. High ketone levels ≥ 100 metabolic substances, nor with reducing metabolites of drugs (e.g. salicylates and Glucose: The reagent area does not react with lactose, galactose, fructose or other

colors on the color chart. Large concentrations of ascorbic acid may decrease or rifampin that might be mistaken for positive bilirubin. The presence of characterized by color development on the test patch that does not correlate with the bilirubin-derived bile pigments may mask the bilirubin reaction. This phenomenon is investigation. Reactions may occur with urine containing large doses of chlorpromazine positive, indicates an underlying pathological condition and requires further Bilirubin: Bilirubin is absent in normal urine, so any positive result, including a trace

of high pigment, and other substances containing sulfhydryl groups may occasionally give reactions up to and including trace (±). Ketone: The test does not react with acetone or β-hydroxybutyrate. Urine specimens

color charl urine has a pH of 7 or greater, add 0.005 to the specific gravity reading indicated on the results. Results are not affected by non-ionic urine components such as glucose. If the Specific Gravity: Ketoacidosis or protein higher than 300 mg/dL may cause elevated

menstruating females. It has been reported that urine of high pH reduces sensitivity, and for erythrocytes. Positive results with this test are often seen with urine from erythrocytes. To enhance accuracy, separate color scales are provided for hemoglobin nemolyzed erythrocytes." while moderate to high concentration of ascorbic acid may inhibit color formation. Blood: A uniform blue color indicates the presence of myoglobin, hemoglobin or Scattered or compacted blue spots indicate intact

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intact erythrocytes reaction. The test is slightly more sensitive to free hemoglobin and myoglobin than to Microbial peroxidase, associated with urinary tract infection, may cause a false positive

readings are not affected by variations in urinary buffer concentration. reagent will run onto the pH area, causing the pH result to appear artificially low. pH phenomenon known as "runover" may occur, in which the acid buffer from the protein pH: If the procedure is not followed and excess urine remains on the strip,

specimens with high specific gravity may give false negative results cleansers containing chlorhexidine may produce false positive results.8 The urine Contamination of urine specimens with quaternary ammonium compounds or skin False positive results may be obtained with highly buffered or alkaline urine mucoprotein.⁸ A negative result does not rule out the presence of these other proteins. highly sensitive for albumin, and less sensitive to hemoglobin, globulin and Protein: Any green color indicates the presence of protein in the urine. This test is

Nitrite: The test is specific for nitrite and will not react with any other substance be obtained if formalin is present. The test cannot be used to detect porphobilinogen. reagent, such as p-aminosalicylic acid and sulfonamides.9 False negative results may normal. A negative result does not at any time preclude the absence of urobilinogen. Urobilinogen: All results lower than 1 mg/dL urobilinogen should be interpreted as The reagent area may react with interfering substances known to react with Ehrlich's

edges should not be interpreted as a positive result. Comparing the reacted reagent

area on a white background may aid in the detection of low nitrite levels, which might proportional to the number of bacteria present in the urine specimen. Pink spots or pink interpreted as a positive result, suggesting the presence of nitrite. Color intensity is not

normally excreted in urine. Any degree of uniform pink to red color should be

when receiving antibiotic therapy or when dietary nitrate is absent. for a sufficient length of time (at least 4 hours) for reduction of nitrate to nitrite to occur. reductase to convert nitrate to nitrite; when urine has not been retained in the bladder negative result does not at any time preclude the possibility of bacteruria. Negative urine specimens with highly buffered alkaline urine or with high specific gravity. A containing less than 0.05 mg/dL nitrite ions. The sensitivity of this test is reduced for otherwise be missed. Ascorbic acid above 30 mg/dL may cause false negatives in urine results may occur in urinary tract infections from organisms that do not contain

bacteria common in urine. color development. The intensity of the color that develops is proportional to the diminish the intensity of the reaction color. This test will not react with erythrocytes or high levels of the drug may cause a false negative reaction. High urinary protein may presence of cephalexin, cephalothin, or high concentrations of oxalic acid may also glucose concentrations (≥ 2,000 mg/dL) may cause test results to be artificially low. The number of leukocytes present in the urine specimen. High specific gravity or elevated cause test results to be artificially low. Tetracycline may cause decreased reactivity, and Leukocytes: The result should be read between 60-120 seconds to allow for complete

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