

Quantimetrix® Dip&Spin®

Urinalysis Dipstick & Microscopics Control Level 1 & 2

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English

Intended Use

The Quantimetrix Dip&Spin Urinalysis Dipstick & Microscopics Control is intended as a control for urinalysis reagent strips, microalbumin, and creatinine by the listed test methods, and as a control for confirmatory tests such as **K-CHECK** and **Ictotest®** reagent tablets, and for **hCG** methods.

In addition, the Dip&Spin Control is intended as a means of validating the processing and centrifugation of patient urine samples prior to the microscopic evaluation of urine sediment.

Summary and Explanation

Control materials having known component concentrations are an integral part of diagnostic procedures. Daily monitoring of control values establishes intralaboratory parameters for accuracy and precision of the test method.

Microscopic QC controls must be run each day the test is performed.¹ Standardized microscopic evaluation of urine sediment is an important part of routine analysis of urine. Along with physical and chemical analysis, microscopic examination of urine can provide valuable information regarding not only renal and urinary tract disease, but also metabolic diseases unrelated to the kidney.² Urinary sediment microscopy generally includes

the detection and identification of red blood cells, leukocytes, epithelial cells, bacteria, casts, and crystals.^{3,4}

Product Description

The Quantimetrix Dip&Spin Urinalysis Dipstick & Microscopics Controls are supplied liquid, ready-to-use in two levels. They do not require reconstitution or dilution. They are prepared from human urine to which stabilized human red and white blood cells, calcium oxalate crystals, and other compounds have been added to produce the desired reactions when tested by the methods indicated in the **Intended Use** section. Preservatives have been added to inhibit microbial growth.

Caution

Contains human urine, human blood cells and human Chorionic Gonadotropin (hCG) from pregnancy urine. The human hCG source material and all blood donor units comprising the human cell source material used in the manufacture of this product have been tested and found nonreactive for Hepatitis B Surface Antigen and Hepatitis C and HIV 1 & 2 antibody when tested by FDA accepted methods. No known test method can assure that a product derived from human material does not contain Hepatitis or HIV virus. Handle the QC material as you would a patient sample. QC materials should be used and disposed of in accordance with regulatory and accreditation requirements.

Warning ⚠ Hazard (H) and Precautionary (P) Statements

Contains Mixture, 3(2H)-isothiazolone, 5-chloro-2-methyl- with 2-methyl-3(2H)-isothiazolone, 1,2-Propylene Glycol, Level 1; 2,4-Pentanedione, Level 2.
H317 – May cause an allergic skin reaction.
P261 – Avoid breathing vapors, mist, or spray.
P272 – Contaminated work clothing should not be allowed out of the workplace.
P280 – Wear protective gloves, protective clothing, and eye protection.
P302+P352 – IF ON SKIN: Wash with plenty of water.
P333+P313 – If skin irritation or rash occurs: Get medical advice/attention.
P362+P364 – Take off contaminated clothing and wash it before reuse.
P501 – Dispose of contents/container in accordance with local, regional, national, and international regulations.
Safety Data Sheet (SDS) available for professional users at quantimetrix.com.

Storage and Stability

The Dip&Spin Control Kit should be at 2°C–8°C when not in use. **Do not freeze.** When stored at 2°C–8°C the controls are stable until the expiration date stated on the label. After opening, the controls will remain stable until the expiration date stated on the label when stored at 2°C–8°C between uses. Discard the control if it becomes more turbid or develops a stronger odor. Discard controls in the same manner as other biological specimens, according to local guidelines.

Procedure for Dipstick Urinalysis and Microscopic Evaluation of Urine Sediment

Remove the controls from the refrigerator and replace the cap on the control bottle with the spout cap included in the control box. Allow the control to come to room temperature (18°C–25°C) for approximately 15–90 minutes depending on the volume remaining in the bottle. Mix the controls thoroughly by inverting the bottle at least 20 times to assure homogeneity of the contents. Avoid foaming. Thorough mixing with each use is important in order to obtain reproducible results. Pour 12 mL of the controls into a standard 15 mL centrifuge tube.

For urinalysis, microalbumin and creatinine testing, immerse the reagent strip in the centrifuge tubes containing the control as if they were patient specimens. Read the urinalysis reagent strips, visually or with an instrumental reader, in accordance with the manufacturer's instructions.

For microscopic evaluation of urine sediment, treat the controls as you would patient samples in accordance with the manufacturer's instructions for the standardized microscopic urinalysis system you are using. The National Committee for Clinical Laboratory Standards (NCCLS) recommends the use of standardized systems in order to yield standardized, reproducible results and to enable the reporting of abnormal sediment elements per unit volume.¹

Procedure for hCG Tests and Confirmatory Tests

Note: The bottles of Level 1 Control are to be used as negative controls for hCG methods. The bottles of Level 2 Control are to be used as positive controls for hCG methods.

Most manufacturers of pregnancy test kits specify the volume of sample to be used with their kits. Many kits include transfer pipets to be used to deliver a certain sample volume onto the test device. It is important that sufficient volume be used to produce the correct test result.

If dispensing the control for hCG tests and confirmatory tests directly from the control bottles, each user should validate that the volume (number of drops) dispensed by the included spout cap is sufficient to meet the pregnancy test kit's and confirmatory tests' requirement for sample volume.

Remove the controls from the refrigerator. Allow the controls to come to room temperature (18°C–25°C) for approximately 15–90 minutes depending on the volume remaining in the bottle. Mix the control thoroughly by inverting the bottle at least 20 times to assure homogeneity of the contents. Avoid foaming. Use the negative and positive controls as if they were patient specimens in accordance with the test kit manufacturer's instructions. If using the same bottle of control dispensed for urinalysis testing and microscopic evaluation, remove the volume of sample to be used for hCG tests and confirmatory tests after centrifugation, before discarding the supernatant and without disturbing the sediment. Immediately close the spout cap and store the controls at 2°C–8°C when not in use.

Expected Values

For visual readings, the expected ranges have been established from interlaboratory data by comparing the dipstick reaction that occurs with the controls to the color comparison chart with multiple lots of each manufacturer's dipsticks or reagent tablets. For expected values for urinalysis reagent strips not listed, please contact Quantimetrix Technical Services.

For instrument readings, the expected ranges have been established from interlaboratory data from multiple lots of each manufacturer's dipsticks. Each laboratory should establish its own precision parameters.

For specific gravity, the expected ranges by refractometer have been established from interlaboratory data.

For hCG, the positive and negative results were obtained by testing each lot number of the controls with multiple lot numbers of different hCG test kits with sensitivities of ≥ 25 mIU/mL.

For microscopic evaluation of urine sediment, the expected ranges for each type of formed element were determined by assay of multiple bottles of the indicated lot by the methods listed. A 12 mL sample volume of the samples were centrifuged at

400 RCF (relative centrifugal force) for 5 minutes. After centrifugation, urine sediment was resuspended in either –0.5 or –1.0 mL of remaining supernatant according to the plasticware manufacturer's directions. The ranges listed are based on the range of elements observed in 10 high power fields. Use of other systems or protocols may yield differing results. Each laboratory should establish its own precision parameters.

Limitations

Any future changes made by the manufacturer of a test method may give different values from the indicated range. Detailed information on the limitations of each test method is included in the limitations section of the manufacturers' package insert. Technical updates can be found on our website. The Quality Control Log can be downloaded from the Quantimetrix website at quantimetrix.com or contact Tech Support at (310) 536-0006, option 3.

Chemstrip/CombiScreen/Combur/Multistix/Urocheck Users

Colors produced by the urobilinogen and/or bilirubin reactions on these dipsticks with the Urinalysis Dipstick Control may not be characteristic of those shown on the manufacturer's label when reading the dipstick reactions visually. The urobilinogen reactions are consistent and intensify with the increase in the urobilinogen concentration but may not provide an exact color match to those displayed on the label.

Note: Siemens® CLINTEK 50 and Siemens® STATUS or CLINTEK STATUS PLUS may see an Albumin/Creatinine ratio result of "Abnormal" with the Level 1 control.

Deutsch

Verwendungszweck

Die Quantimetrix Dip&Spin Urinalysis Dipstick & Microscopics Control ist als Kontrolle für Urinalyse-Reagenzstreifen, Mikroalbumin und Kreatinin gemäß den aufgeführten Testmethoden sowie als Kontrolle für Bestätigungstests wie z. B. **K-CHECK** und **Ictotest®** Reagenz-Tabletten und für **hCG**-Methoden bestimmt.

Darüber hinaus dient die Dip&Spin Control zur Bewertung der Verarbeitung und Zentrifugierung von Patienten-Urinproben vor der mikroskopischen Beurteilung des Urinsediments.

Zusammenfassung und Erklärung

Kontrollmaterialien mit bekannten Konzentrationen von Komponenten sind ein integraler Bestandteil diagnostischer Verfahren. Im Rahmen der täglichen Überwachung von Kontrollwerten werden laborinterne Parameter für die Genauigkeit und Präzision der Testmethode festgelegt.

An jedem Tag, an dem der Test durchgeführt wird, müssen mikroskopische Qualitätskontrollen (QC) laufen.¹ Die standardisierte mikroskopische Beurteilung von Urinsediment ist ein wichtiger Bestandteil der routinemäßigen Urinalyse. Zusammen mit der physikalischen und chemischen Analyse kann die mikroskopische Untersuchung des Urins wertvolle Informationen nicht nur über Erkrankungen von Nieren und Harnwegen, sondern auch über von der Niere unabhängige Stoffwechselerkrankungen liefern.² Zur mikroskopischen Untersuchung von Urinsediment gehört grundsätzlich der Nachweis und die Identifizierung von roten Blutkörperchen, Leukozyten, Epithelzellen, Bakterien, Ausgüssen und Kristallen.^{3,4}

Produktbeschreibung

Die Quantimetrix Dip&Spin Urinalysis Dipstick & Microscopics Controls werden gebrauchsfertig in zwei Stufen ausgeliefert. Es ist keine Rekonstitution oder Verdünnung erforderlich. Sie werden aus menschlichem Urin hergestellt, der mit stabilisierten roten und weißen Blutkörperchen, Kalziumoxalatkristallen und anderen Substanzen angereichert wurde, um die gewünschte Reaktion zu erzeugen, wenn das Produkt gemäß den unter **Verwendungszweck** beschriebenen Verfahren eingesetzt wird. Das Produkt wurde mit Konservierungsstoffen angereichert, um mikrobiellem Keimwachstum entgegenzuwirken.

Warnhinweis

Enthält menschliches Urin, menschliche Blutkörperchen und menschliches Choriongonadotropin (hCG) aus Urin bei Schwangerschaft. Das menschliche hCG-Quellenmaterial und alle bei der Produktherstellung verwendeten Blutspenden, die das menschliche Zellenquellenmaterial beinhalten, wurden unter Einhaltung anerkannter FDA-Methoden auf Hepatitis B-Oberflächenantigene, Hepatitis C und Antikörper gegen HIV 1 & 2 getestet. Die Testergebnisse waren nicht-reaktiv. Es sind keine Testmethoden bekannt, mit denen garantiert werden kann, dass die aus menschlichem Material gewonnenen Produkte frei von Hepatitis- oder HIV-Viren sind. Die Materialien für die Qualitätskontrolle sollten wie Patientenproben gehandhabt werden. Die Materialien müssen im Einklang mit den gesetzlichen Bestimmungen und Zulassungsvorschriften verwendet und entsorgt werden.

Achtung ⚠ Gefahrenhinweise (H) Sicherheitshinweise (P)

Gemisch, 3(2H)-isothiazolone, 5-chloro-2-methyl- mit 2-methyl-3(2H)-isothiazolone, 1,2-Propylene Glycol, Stufe-1; 2,4-Pentanedione, Stufe-2.
H317 – Kann allergische Hautreaktionen verursachen.
P261 – Einatmen von Nebel, Dämpfen, Aerosol vermeiden.
P272 – Kontaminierte Arbeitskleidung nicht außerhalb des Arbeitsplatzes tragen.
P280 – Schutzhandschuhe, Schutzkleidung und Augenschutz tragen.
P302+P352 – BEI KONTAKT MIT DER HAUT: Mit viel Wasser waschen.
P333+P313 – Bei Hautreizung oder -ausschlag: Ärztlichen Rat einholen/ärztliche Hilfe hinzuziehen.
P362+P364 – Alle kontaminierten Kleidungsstücke sofort ausziehen und vor erneutem Tragen waschen.
P501 – Inhalt/Behälter entsprechend örtlichen, regionalen, nationalen und internationalen Richtlinien der Entsorgung zuführen. Sicherheitsdatenblatt (SDS) steht Ihnen im Internet unter quantimetrix.com zur Verfügung.

Lagerung und Stabilität

Das Dip&Spin Control Kit sollte bei Nichtgebrauch bei 2°C bis 8°C gelagert werden. **Nicht einfrieren.** Bei Lagerung bei 2°C bis 8°C sind die Kontrollen bis zum auf dem Etikett angegebenen Verfallsdatum stabil. Nach dem Öffnen bleiben die Kontrollen bis zum auf dem Etikett angegebenen Verfallsdatum stabil, wenn sie zwischen den Verwendungen bei 2°C bis 8°C gelagert werden. Falls die Kontrolle trüb wird oder einen starken Geruch ausstrahlt, sollte sie entsorgt werden. Kontrollen auf gleiche Weise wie andere biologische Proben gemäß den örtlichen Richtlinien entsorgen.

Verfahren für Dipstick Urinalyse und mikroskopische Beurteilung von Urinsediment

Nehmen Sie die Kontrollen aus dem Kühlschrank, und tauschen Sie die Kappe des Kontrollfläschchens gegen den in der Kontrollbox enthaltenen Ausgießverschluss aus. Lassen Sie die Kontrollen je nach der noch im Fläschchen verbliebenen Menge ca. 15 bis 90 Minuten lang auf Raumtemperatur (18°C bis 25°C) aufwärmen. Mischen Sie die Kontrollen gründlich, indem Sie das Fläschchen mindestens 20 Mal umdrehen und so einen homogenen Inhalt sicherstellen. Nicht schäumen lassen. Ein gründliches Mischen vor jeder Verwendung ist unerlässlich, um reproduzierbare Resultate zu erhalten. Gießen Sie

12 ml der Kontrollen in ein standardmäßiges 15 ml Zentrifugenröhrchen.

Zur Urinalyse, Mikroalbumin- und Creatinin-Testung tauchen Sie den Reagenzstreifen wie bei einer Patientenprobe in das Zentrifugenröhrchen mit der Kontrolle. Die Urinalyse-Teststreifen visuell oder in einem Lesegerät gemäß den Herstelleranweisungen ablesen.

Zur mikroskopischen Beurteilung von Urinsediment behandeln Sie die Kontrollen wie Patientenproben entsprechend den Herstelleranweisungen für das von Ihnen verwendete, standardisierte, mikroskopische Urinalysesystem. Das National Committee for Clinical Laboratory Standards (NCCLS) empfiehlt die Verwendung standardisierter Systeme, um standardisierte, reproduzierbare Ergebnisse zu erhalten und die Angabe abnormaler Sedimentbestandteile pro Volumeneinheit zu ermöglichen.¹

Verfahren für hCG-Tests und Bestätigungstests

Hinweis: Die Fläschchen mit der Level-1-Kontrolle sind bei hCG-Methoden als negative Kontrollen vorgesehen. Die Fläschchen mit der Level-2-Kontrolle sind bei hCG-Methoden als positive Kontrollen vorgesehen.

Die meisten Hersteller von Schwangerschaftstests geben die für ihre Tests benötigte Probenmenge an. Viele Testkits enthalten Pipetten zum Übertragen einer bestimmten Probenmenge auf das Testgerät. Es ist wichtig, dass eine ausreichende Probenmenge verwendet wird, um das richtige Testergebnis zu erzielen.

Falls die Kontrolle für die hCG-Tests und die Bestätigungstests direkt von den Kontrollfläschchen aus verabreicht wird, muss der Benutzer bestätigen, dass die über den Tropfverschluss verabreichte Menge (Anzahl der Tropfen) ausreicht, damit sie die Anforderungen an die Probenmenge für den Schwangerschaftstest und für die Bestätigungstests erfüllt.

Die Kontrollen aus dem Kühlschrank nehmen. Die Kontrollen ca. 15 bis 90 Minuten lang auf Raumtemperatur (18°C bis 25°C) aufwärmen lassen, je nach der noch im Fläschchen verbliebenen Menge. Mischen Sie die Kontrollen gründlich, indem Sie das Fläschchen mindestens 20 Mal umdrehen und so einen homogenen Inhalt sicherstellen. Nicht schäumen lassen. Verwenden Sie die negativen und positiven Kontrollen entsprechend den Herstelleranweisungen des Testkits wie Patienten-Proben. Bei Verwendung der gleichen Flasche der Kontrolle für die Urinalyse und die mikroskopische Auswertung sollte das für die hCG-Tests bzw. die Bestätigungstests verwendete Probenvolumen nach der Zentrifugierung entfernt werden, bevor der Überstand entsorgt wird. Die Sedimente dürfen dabei nicht aufgeschüttelt werden. Ausgießverschluss sofort verschließen und Kontrollen bei Nichtgebrauch bei 2°C bis 8°C lagern.

Erwartete Werte

Für visuelle Messungen wurden die erwarteten Bereiche aus den Daten verschiedener Labors bestimmt, indem die mit den Kontrollen erhaltene Teststäbchenreaktion mit der Farbvergleichstabelle verglichen wurde, die Farben für mehrere Chargen der Teststäbchen bzw. Reagenztabletten jedes Herstellers enthält. Erwartete Werte für nicht aufgeführte Urinalyse-Reagenzstreifen sind von Quantimetrix Technical Services erhältlich.

Für Gerätemessungen wurden die erwarteten Werte anhand von Daten verschiedener Labors und mehreren Chargen von Teststäbchen jedes Herstellers bestimmt. Jedes Labor sollte seine eigenen Präzisionsparameter bestimmen.

Für relative Dichte wurden die mit dem Refraktometer ermittelten, erwarteten Bereiche aus Daten von verschiedenen Labors bestimmt.

l'omogeneidad del contenido. Evitar la formación de espuma. Utilizar los controles negativos y positivos como se si tratase del campione del paciente, en base a las instrucciones del fabricante del kit de test. Si se usa el mismo frasco del control ergo para los análisis de las orinas y para la valoración microscópica, remover el volumen del campione a usarse para i test hCG e i confirma dopo la centrifugazione, prima di eliminare il supernatante e senza disturbare il sedimento. Chiedere immediatamente il beccuccio e conservare i controlli a temperatura comprese tra 2°C-8°C fra i vari impieghi.

Valori previsti

Per le lettura visuali, i range previsti sono stati stabiliti attraverso dati di diversi laboratori, confrontando la reazione dei dipstick con quelli della carta dei colori, utilizzando diversi lotti di ogni dipstick o di pastiglia reagente dei vari fabbricanti. Per conoscere i valori previsti delle strisce per l'esame delle urine non in elenco, contattare il servizio tecnico Quantimetrix.

In relazione alle lettura con appositi strumenti i range previsti sono stati stabiliti da dati di vari laboratori su diversi lotti di ogni fabbricante di dipstick. Ogni laboratorio dovrà stabilire i propri parametri di precisione.

In relazione alla gravità specifica, gli ambiti previsti con l'uso del rifrattometro sono stati stabiliti attraverso i dati di diversi laboratori.

In relazione a hCG, i risultati positivi e negativi sono stati ottenuti testando ogni numero di lotto dei controlli con molteplici numeri di lotto di diversi kit di test hCG con sensibilità di ± 25 mIU/mL.

In relazione alla valutazione microscopica del sedimento urinario, gli ambiti previsti per ogni tipo di elemento formato è stato determinato mediante analisi di molteplici lotti del lotto indicato, utilizzando il metodo elencato. 12 ml di campione sono stati centrifugati a 400 RCF (forza centrifuga relativa) per 5 minuti. Dopo la centrifugazione il sedimento urinario è stato riscoperto in ~0,5 o ~1,0 ml del supernatante restante, in base alle istruzioni del fabbricante dell'articolo di plastica. Gli ambiti elencati fanno riferimento al range degli elementi osservati in 10 campi ad alto ingrandimento. L'impiego di altri sistemi o protocolli può portare a risultati differenti. Ogni laboratorio dovrà stabilire i suoi propri parametri di precisione.

Limiti

Eventuali futuri cambiamenti apportati dal fabbricante di un metodo di analisi potrebbero dare valori diversi dall'intervallo di valori indicato. Informazioni dettagliate sui limiti di ciascun metodo di analisi sono incluse nella sezione Limiti dell'insero informativo del fabbricante. Aggiornamenti tecnici sono reperibili sul nostro sito web. Il registro del controllo della qualità si può ottenere scaricandolo dal sito web Quantimetrix all'indirizzo quantimetrix.com oppure contattando il team del Supporto tecnico al numero +1 (310) 536-0006, opzione 3.

Utilizzatori di Chemstrip/CombiScreen/Combur/Multistix/Urocheck

I colori prodotti dalle reazioni di urobilinogeno e/o bilirubina su questi dipstick con il Controllo dipstick urina potrebbero non rispecchiare quelli illustrati sull'etichetta del fabbricante quando le reazioni dei dipstick vengono lette visivamente. Le reazioni dell'urobilinogeno sono costanti e aumentano di intensità all'aumentare della concentrazione di urobilinogeno ma è possibile che non vi sia un'esatta corrispondenza di colore con quelle mostrate sull'etichetta.

Nota: Siemens® CLINITEK 50 e Siemens® STATUS o CLINITEK STATUS PLUS potrebbero riscontrare risultati "Anormali" per il rapporto albumina/creatinina con il controllo di Livello 1.

Español

Uso previsto

El Quantimetrix Dip&Spin Urinalysis Dipstick & Microscopics Control se utiliza como control para las tiras reactivas de análisis de orina, microalbumina y creatinina por los métodos indicados, y como control de pruebas de confirmación como las tabletas reactivas **K-CHECK** e **Ictest**® y para los métodos de detección de **hCG**.

Además, el Dip&Spin Control se utiliza para validar el procesado y centrifugado de muestras de orina de pacientes antes de la evaluación microscópica de la sedimentación presente en la orina.

Resumen y explicación

Los materiales de control que tienen concentraciones conocidas del componente forman parte integral de los procedimientos diagnósticos. La monitorización diaria de los valores de control establece los parámetros de exactitud y precisión del método de análisis en cada laboratorio.

Los controles microscópicos de control de calidad deben realizarse cada día que se lleva a cabo la prueba.¹ La evaluación microscópica normalizada de la sedimentación presente en la orina es una parte importante del análisis rutinario de la orina. Junto con el análisis físico y químico, el estudio microscópico de la orina puede aportar valiosa información no sólo sobre enfermedades renales y del tracto urinario, sino también sobre enfermedades metabólicas que no tengan relación alguna con el riñón.² El estudio microscópico de la sedimentación presente en la orina generalmente incluye la detección e identificación de hemáticas, leucocitos, células epiteliales, bacterias, cilindros y cristales.^{3,4}

Descripción del producto

Los Quantimetrix Dip&Spin Urinalysis Dipstick & Microscopics Controls se suministran líquidos, listos para usar en dos niveles. No requieren reconstitución o dilución. Están preparados a partir de orina humana a la que se han agregado glóbulos humanos rojos y blancos estandarizados, cristales de oxalato de calcio y otros compuestos para producir las reacciones deseadas cuando se prueban con los métodos indicados en la sección **Uso previsto**. Se han agregado conservantes para inhibir la proliferación microbiana.

Precaución

Contiene orina humana, células sanguíneas humanas y gonadotropina coriónica humana (hCG) de la orina del embarazo. El material fuente del hCG humano y de todas las unidades donantes de sangre que comprenden el material fuente de células humanas utilizado en la fabricación de este producto se ha probado y no se ha detectado ningún reactivo para el antígeno de superficie de la Hepatitis B ni anticuerpos de Hepatitis C y VIH 1 y 2 cuando las pruebas se realizan con métodos aceptados por la FDA. Ningún método de prueba conocido puede asegurar que un producto derivado de material humano no contenga hepatitis o virus VIH. Trabaje con el material QC como lo haría con una muestra de paciente. Los materiales QC deben usarse y eliminarse de acuerdo con los requisitos regulatorios y de acreditación.

Atención ⚠️ Indicaciones de peligro (H) Consejos de precaución (P)

Mezcla, 3(2H)-isothiazolone, 5-chloro-2-methyl- con 2-methyl-3(2H)-isothiazolone,

1,2-Propylene Glycol, nivel 1; 2,4-Pentanedione, nivel 2.

H317 – Puede causar una reacción alérgica cutánea.

P261 – Evite respirar vapores, niebla o aerosol.

P272 – La ropa de trabajo contaminada no debe sacarse del lugar de trabajo.

P280 – Lleve guantes, prendas y gafas de protección.

P302+P352 – EN CASO DE CONTACTO CON LA PIEL: lave con agua abundante.

P333+P313 – Si aparece irritación o erupción cutánea: consulte a un médico.

P362+P364 – Quite se la ropa contaminada y lávela antes de volver a utilizarla.

P501 – Elimine el contenido/contenedor conforme a la normativa local, regional, nacional e internacional vigente.

La hoja de datos de seguridad (SDS) está disponible para los usuarios profesionales en quantimetrix.com.

Almacenamiento y estabilidad

El Dip&Spin Control Kit deberá almacenarse a 2°C-8°C cuando no se utilice. **No congelar**. Cuando se almacenan a 2°C-8°C, los controles permanecen estables hasta la fecha de caducidad que figura en la etiqueta. Una vez abiertos, los controles permanecerán estables hasta la fecha de caducidad que figura en la etiqueta cuando se almacenen a 2°C-8°C después de cada uso. Deseche el control si se vuelve más turbio o si desarrolla un olor más fuerte. Desechar los controles de la misma forma que cualquier otra muestra biológica, conforme a las normativas locales.

Procedimiento para el análisis de orina con tira reactiva y la evaluación microscópica de la sedimentación presente en la orina

Extraiga los controles de la nevera y sustituya la tapa del frasco de control por la tapa del surtidor incluida en la caja de control. Deje que el control se estabilice a temperatura ambiente (18 °C-25 °C) durante aproximadamente 15-90 minutos, dependiendo del volumen que quede en el frasco. Mezcle bien los controles invirtiendo el frasco por lo menos 20 veces para garantizar la homogeneidad del contenido. Evite la formación de espuma. Para poder obtener resultados reproducibles, es importante mezclar bien los controles cada vez que se utilicen. Vierta 12 ml de los controles en un tubo de centrifuga estándar de 15 ml.

Para los análisis de orina, microalbuminuria y creatinina, sumerja la tira reactiva en los tubos de centrifuga que contienen el control, igual que si fueran muestras de pacientes. Lea las tiras reactivas de análisis de orina, visualmente o con un instrumento lector, de acuerdo con las instrucciones del fabricante.

Para la evaluación microscópica de la sedimentación presente en la orina, los controles deberán tratarse como si fueran muestras de pacientes, de acuerdo con las instrucciones del fabricante para el sistema microscópico normalizado de análisis de orina que esté utilizando. El National Committee for Clinical Laboratory Standards (NCCLS) recomienda el empleo de sistemas normalizados con el fin de obtener resultados reproducibles y normalizados, y poder detectar e informar acerca de la presencia de elementos anormales en la sedimentación en cada volumen unitario.¹

Procedimiento para los ensayos de hCG y los ensayos de confirmación

Nota: Los frascos de control de concentración 1 se deben usar como controles negativos de los métodos de hCG. Los frascos de control de concentración 2 se deben usar como controles positivos de los métodos de hCG.

La mayoría de los fabricantes de kits de prueba de embarazo especifican el volumen de muestra a utilizar en sus kits. Muchos kits incluyen pipetas de transferencia que se utilizan para suministrar un determinado volumen de muestra en el dispositivo de prueba. Es importante que se use suficiente volumen para producir el resultado correcto de la prueba.

Si el control para pruebas hCG y pruebas confirmatorias se dispensa directamente desde de los frascos de los controles, cada usuario debe validar que el volumen (cantidad de gotas) dispensado por la tapa del surtidor sea suficiente para cumplir con los requisitos del kit de prueba de embarazo y las pruebas confirmatorias.

Extraiga los controles de la nevera. Deje que los controles se estabilicen a temperatura ambiente (18 °C-25 °C) durante aproximadamente 15-90 minutos, dependiendo del volumen que quede en el frasco. Mezcle bien los controles invirtiendo el frasco por lo menos 20 veces para garantizar la homogeneidad del contenido. Evite la formación de espuma. Use los controles positivo y negativo como si fueran muestras de paciente, de acuerdo con las instrucciones del fabricante del kit de análisis. Si el mismo frasco de control dispensado se utiliza para las pruebas de análisis de orina y la evaluación microscópica, retire el volumen de muestra a utilizar en las pruebas de hCG y las pruebas de confirmación después del centrifugado, antes de descartar el sobrenadante y sin perturbar el sedimento. Cierre inmediatamente la tapa del surtidor y almacene los controles a 2 °C-8 °C cuando no se utilicen.

Valores esperados

En el caso de lecturas visuales, los intervalos esperados se han establecido a partir de datos de varios laboratorios, comparando la reacción de la tira reactiva que se produce con los controles, con la carta de comparación de colores de varios lotes de tiras reactivas o tabletas de reactivo de cada fabricante. En cuanto a los valores esperados de las tiras de reactivo para análisis de orina que no figuren, póngase en contacto con el Servicio Técnico de Quantimetrix.

En el caso de lecturas con instrumento, los intervalos esperados se han establecido a partir de datos obtenidos en varios laboratorios con múltiples lotes de tiras reactivas de cada fabricante. Cada laboratorio deberá establecer sus propios parámetros de precisión.

En el caso del peso específico, los intervalos esperados con el refractómetro se han establecido a partir de datos obtenidos en varios laboratorios.

En el caso de hCG, los resultados positivo y negativo se obtuvieron analizando cada número de lote de los controles con múltiples números de lote de diferentes kits de análisis de hCG con sensibilidades de ≥ 25 mIU/mL.

En el caso de la evaluación microscópica de la sedimentación presente en la orina, los intervalos esperados para cada tipo de elemento formado se determinaron mediante valoración de varios frascos del lote indicado por medio de los métodos listados. Se centrifugó un volumen de las muestras de 12 ml a 400 RCF (fuerza centrifuga relativa) durante 5 minutos. Tras la centrifugación, la sedimentación presente en la orina se volvió a suspender en ~0,5 o ~1,0 ml del sobrenadante restante, de acuerdo con las instrucciones del fabricante de los plásticos. Los intervalos listados se basan en el intervalo de elementos observados en 10 campos de gran aumento. El uso de otros sistemas o protocolos puede arrojar resultados distintos. Cada laboratorio deberá establecer sus propios parámetros de precisión.

Limitaciones

Cualquier cambio futuro realizado por el fabricante de un método de prueba puede dar valores diferentes del rango indicado. En la sección de limitaciones del prospecto del fabricante se incluye información detallada sobre las limitaciones de cada método de prueba. En nuestro sitio web se pueden encontrar las actualizaciones técnicas. El registro de control de calidad se puede descargar en el sitio web de Quantimetrix en quantimetrix.com poniéndose en contacto con el Soporte técnico en el +1 (310) 536-0006, opción 3.

Usuarios de Chemstrip/CombiScreen/Combur/Multistix/Urocheck

Los colores producidos por las reacciones al urobilinogeno y/o a la bilirubina en esas tiras reactivas con el Control de tiras reactivas en orina podrían no ser características de las que se indican en la etiqueta del fabricante al leer visualmente las reacciones en la tira reactiva. Las reacciones de urobilinogeno son coherentes y se intensifican cuando aumenta la concentración de urobilinogeno, pero puede que no den colores exactamente iguales a los que se muestran en la etiqueta.

Nota: Siemens® CLINITEK 50 y Siemens® STATUS o CLINITEK STATUS PLUS pueden ver un resultado en la proporción de albumina/creatinina calificado de "Anormal" con el control de Nivel 1.

References | Bibliographie | Références | Bibliografia | Bibliografía

1 Routine Urinalysis and Collection, Transportation, and Preservation of Urine Specimens, NCCLS Documentation GP 16-A, Approved Guideline, (1995), NCCLS, Wayne PA, 19087.

2 Bradley GM, Benson ES, Todd-Sanford Clinical Diagnosis by Laboratory Methods, 15th ed., Philadelphia, PA: Saunders, 1974.

3 Bologna, CV, Understanding Laboratory Medicine, St. Louis, MO: CV Mosby Co., 1971.

4 Dudas HC, Lab Med, 12:765, 1981.

Analyte/Method	Level 1 - 55251	Level 2 - 55252	Units
Red Blood Cells (Erythrocytes)			
Beckman Coulter IRIS Diagnostics IQ®200	0 - 10	8 - 43	/ μ L
Series Analyzers			
KOVA® GLASSTIC® SLIDE 10 with GRIDS	3 - 13	18 - 95	/ μ L
Non-grid slides (~0.5 mL)	0 - 5	4 - 38	/hpf
Non-grid slides (~1.0 mL)	0 - 4	1 - 25	/hpf
Ken-Slide	0 - 5	3 - 40	/hpf
Slide & Coverslip (~0.5 mL)	0 - 6	3 - 21	/hpf
Slide & Coverslip (~1.0 mL)	0 - 4	1 - 13	/hpf
FisherBrand UriSystem DeciSlide	0 - 5	4 - 31	/hpf
White Blood Cells (Leukocytes)			
Beckman Coulter IRIS Diagnostics IQ®200	2 - 15	25 - 71	/ μ L
Series Analyzers			
KOVA® GLASSTIC® SLIDE 10 with GRIDS	2 - 15	17 - 85	/ μ L
Non-grid slides (~0.5 mL)	0 - 9	1 - 34	/hpf
Non-grid slides (~1.0 mL)	0 - 4	1 - 19	/hpf
Ken-Slide	0 - 6	0 - 41	/hpf
Slide & Coverslip (~0.5 mL)	0 - 4	1 - 19	/hpf
Slide & Coverslip (~1.0 mL)	0 - 4	0 - 10	/hpf
FisherBrand UriSystem DeciSlide	0 - 5	2 - 22	/hpf
Casts			
Beckman Coulter IRIS Diagnostics IQ®200	None	None	
Series Analyzers			
KOVA® GLASSTIC® SLIDE 10 with GRIDS	None	None	
Non-grid slides (~0.5 mL)	None	None	
Non-grid slides (~1.0 mL)	None	None	
Ken-Slide	None	None	
Slide & Coverslip (~0.5 mL)	None	None	
Slide & Coverslip (~1.0 mL)	None	None	
Crystals (calcium oxalate dihydrate with some monohydrate and amorphous forms)			
Beckman Coulter IRIS Diagnostics IQ®200	None	Present	
Series Analyzers			
KOVA® GLASSTIC® SLIDE 10 with GRIDS	none	present	
Non-grid slides (~0.5 mL)	none	present	
Non-grid slides (~1.0 mL)	none	present	
Ken-Slide	none	present	
Slide & Coverslip (~0.5 mL)	none	present	
Slide & Coverslip (~1.0 mL)	none	present	
FisherBrand UriSystem DeciSlide	none	present	

Analytes/Method	Level 1 - 55251	Level 2 - 55252	Units
Bacteria			
Beckman Coulter IRIS Diagnostics iQ®200 Series Analyzers	May be present	Present	
KOVA® GLASSTIC® SLIDE 10 with GRIDS	May be present	Present	
Non-grid slides (~0.5 mL)	May be present	Present	
Non-grid slides (~1.0 mL)	May be present	Present	
Gen-Slide	May be present	Present	
Slide & Coverslip (~0.5 mL)	May be present	Present	
Slide & Coverslip (~1.0 mL)	May be present	Present	
FisherBrand UriSystem DeciSlide	May be present	Present	

Analytes	Level 1 - 55251	Level 2 - 55252
Accutest® URS 10 Urine Reagent Strips (VISUAL)		
Leukocytes	Negative	15 - 500 cells/µL (Tr - Lg)
Nitrite	Negative	Positive
Urobilinogen	Normal (0.2 - 1 mg/dL)	2 - 8 mg/dL ⁷
Protein	Negative	30 - 2000 mg/dL (1+ - 4+)
pH	5.0 - 6.0	7.0 - 8.5
Blood	Negative - Trace	25 - 200 cells/µL (Sm - Lg)
Specific Gravity	1.005 - 1.020	1.000 - 1.015
Ketones	Negative	5 - 160 mg/dL (Tr - Lg)
Bilirubin	Negative	Small - Large
Glucose	Negative	100 - 1000 mg/dL (Tr - 3+)

Accutest® (Analyzers) ®		
Glucose	Negative	100 - 1000 mg/dL (Tr - 3+)
Bilirubin	Negative	Small - Large
Ketones	Negative	5 - ≥80 mg/dL (Tr - Lg)
Specific Gravity	1.010 - ≥1.030	1.000 - 1.015
Blood	Negative - Trace	25 - 200 cells/µL (Sm - Lg)
pH	5.0 - 6.5	7.0 - 8.5
Protein	Negative	Trace - 300 mg/dL
Urobilinogen	Normal (0.2 mg/dL)	2 - 8 mg/dL ⁷
Nitrite	Negative	Positive
Leukocytes	Negative	15 - 500 cells/µL (Tr - Lg)

Accustrip® URS Reader/Visual		
Bilirubin	Negative	1 - 4 mg/dL (1+ - 3+) (17 - 70 µmol/L) ⁷
Blood	Negative - 10 Ery/µL	10 - 250 Ery/µL
Glucose	Negative - Normal	50 - ≥500 mg/dL (8.3 - ≥27.8 mmol/L)
Ketones	Negative	25 - 300mg/dL (1+ - 3+) (2.5 - 30mmol/L)
Leukocytes	Negative	25 - 500 Leu/µL
Nitrite	Negative	Positive
pH	5.0 - 7.0	7.0 - 9.0
Protein	Negative	30 - 500 mg/dL (0.3 - 5.0 g/L) (1+ - 3+)
Specific Gravity (Density)	1.010 - 1.025	1.000 - 1.020
Urobilinogen	Normal	2 - 12 mg/dL (34 - 200µmol/L) ⁷

Beckman Coulter IRIS Diagnostics® iChem®VELOCITY™ Analyzer		
Bilirubin	Negative	Not Compatible
Urobilinogen	Normal	2 - 4 mg/dL (1+ - 2+)
Ketones	Negative	5 - 80 mg/dL (Tr+ - 2+)
Ascorbic Acid	Negative	Negative
Glucose	Negative	50 - 500 mg/dL (1+ - 3+)
Protein	Negative	30 - 500 mg/dL (1+ - 3+)
Blood	Negative - Trace	0.03 - ≥1 mg/dL (1+ - 3+)
pH	5.0 - 7.0	7.0 - 9.0
Nitrite	Negative	Positive
Leukocytes	Negative	75 - 500 mg/dL (2+ - 3+)
Specific Gravity	1.020 - 1.026	1.012 - 1.018

Confirmatory and Other Tests		
K-CHECK (Ketones)	Negative	Small - Large
Ictotest (Bilirubin)	Negative	Positive
Refractometer (Specific Gravity)	1.021 - 1.027	1.013 - 1.019
hCG	Negative	Positive
pH Paper	4.0 - 6.0	7.0 - 9.0
Sulfosalicylic Acid (Total Protein)	(≤ 0.05) ¹⁰	Positive (≥0.50) ¹⁰

Analytes	Level 1 - 55251	Level 2 - 55252
Henry Schein One Step Plus/UriSpec Plus Analyzer (Visual)		
Bilirubin	Negative	1 - 4 mg/dL (1+ - 3+) (17 - 70 µmol/L) ⁴
Blood	Negative - 10 Ery/µL	10 - 250 Ery/µL
Glucose	Negative - Normal	50 - ≥500 mg/dL (8.3 - ≥27.8 mmol/L) ⁸
Ketones	Negative	25 - 300 mg/dL (1+ - 3+) (2.5 - 30 mmol/L)
Leukocytes	Negative	25 - 500 Leu/µL
Nitrite	Negative	Positive
pH	5.0 - 7.0	7.0 - 9.0
Protein	Negative	30 - 500 mg/dL (0.3 - 5.0 g/L) (1+ - 3+)
Specific Gravity (Density)	1.005 - 1.025	1.000 - 1.020
Urobilinogen	Normal	2 - 12 mg/dL (34 - 200 µmol/L) ⁴

Henry Schein UriSpec® 10SG (Visual)		
Leukocytes	Negative	15 - 500 Cells/µL (Tr - Lg)
Nitrite	Negative	Positive
Urobilinogen	Normal (0.2 E.U./dL)	1 - 8 E.U./dL
Protein	Negative	Trace - 300 mg/dL (Tr - 3+)
pH	5.0 - 6.5	6.5 - 8.5
Blood	Negative - 10 Ery/µL	10 - 200 Cells/µL (Tr - Lg)
Specific Gravity (1.0-)	1.015 - 1.030	1.000 - 1.015
Ketones	Negative	5 - 160 mg/dL (Tr - Lg)
Bilirubin	Negative	Small - Large (1+ - 3+)
Glucose	Negative	100 - 1000 mg/dL (Tr - 3+)

MACHEREY-NAGEL® URYXXON® Relax/300/500 Analyzer / VISUAL		
Bilirubin	Negative	1 - 4 mg/dL (1+ - 3+) (17 - 70 µmol/L) ⁷
Blood	Negative - Trace	10 - 250 Ery/µL
Glucose	Negative - Normal	50 - ≥500 mg/dL (8.3 - ≥27.8mmol/L) ⁸
Ketones	Negative	25 - 300 mg/dL (1+ - 3+) (2.5 - 30 mmol/L)
Leukocytes	Negative	25 - 500 Leu/µL
Nitrite	Negative	Positive
pH	5.0 - 7.0	7.0 - 9.0
Protein	Negative	30 - 500 mg/dL (0.3 - 5.0 g/L) (1+ - 3+)
Specific Gravity	1.005 - 1.025	1.000 - 1.020
Urobilinogen	Normal	2 - 12 mg/dL (34 - 200 µmol/L) ⁷

McKesson® (Consult Diagnostics 10SG Urine Reagent Strips) (Visual)		
Glucose	Negative	100 - 1000 mg/dL (± - 3+)
Bilirubin	Negative	1 - 4 mg/dL (1+ - 3+)
Ketones	Negative	5 - 160 mg/dL (± - 4+)
Specific Gravity	1.015 - 1.030	1.000 - 1.020
Blood	Negative - Trace	1+ - 3+
pH	5.0 - 6.5	7.0 - 9.0
Protein	Negative	15 - 300 mg/dL (± - 3+)
Urobilinogen	Normal (0.2 E.U./dL)	1 - 8 mg/dL
Nitrite	Negative	Positive
Leukocytes	Negative	15 - 500 Leu/µL (± - 3+)

McKesson® 120 Urine Analyzer		
Leukocytes	Negative	15 - 500 Leu/µL (± - 3+)
Nitrite	Negative	Positive
Urobilinogen	Normal (0.2 E.U./dL)	0.2 - 2.0 mg/dL ¹¹
Protein	Negative	Trace - 100 mg/dL (± - 2+)
pH	5.0 - 6.5	7.0 - 9.0
Blood	Negative - Trace	25 - 200 Ery/µL (1+ - 3+)
Specific Gravity	1.015 - 1.030	1.000 - 1.020
Ketones	Negative	5 - 80 mg/dL (Tr - 3+)
Bilirubin	Negative	1 - 4 mg/dL (1+ - 3+)
Glucose	Negative	100 - ≥1000 mg/dL (Tr - 3+)

ROCHE VISUAL TESTING (Visual Test Strips Only)		
Specific Gravity	1.015 - 1.030	1.000 - 1.010
pH	5 - 6	7 - 9
Leukocytes	Negative	1+ - 2+
Nitrite	Negative	Positive
Protein	Negative	30 - 100 mg/dL
Glucose	Normal	250 - 1000 mg/dL
Ketones	Negative	Small - Large (1+ - 3+)
Urobilinogen ^{9*}	Normal	1 - 8 mg/dL (1+ - 3+)
Bilirubin ^{9*}	Negative	1+ - 3+
Blood	Negative - Trace	50 - 250 Ery/µL
Microalbumin ⁹	Negative - 20 mg/L	50 - 100 mg/L

Analytes	Level 1 - 55251	Level 2 - 55252
ROCHE Chemstrip 101 Urine Analyzer or ROCHE Urisys 1100 Urine Analyzer		
Blood	Negative - Trace	50 - 250 Ery/ μ L (1+ - 2+)
Bilirubin	Negative	3 - 6 mg/dL (2+ - 3+)*
Urobilinogen	Normal	1 - 8 mg/dL (1+ - 3+) ^{7*}
Ketones	Negative	15 - 150 mg/dL (1+ - 3+)
Glucose	Normal	250 - 1000 mg/dL (2+ - 3+)
Protein	Negative ⁶	15 (Tr) - 100 mg/dL (Tr - 2+) ⁷
Nitrite	Negative	Positive
Leukocytes	Negative	75 - 500 Leu/ μ L (1+ - 2+)
pH	5 - 6.5	7 - 9
Specific Gravity	1.015 - 1.025	1.000 - 1.015
ROCHE Chemstrip UA Urine Analyzer		
Blood	Negative - Trace	150 - 250 Ery/ μ L
Bilirubin	Negative	3 - 6 mg/dL (2+ - 3+)*
Urobilinogen	Normal	1 - 8 mg/dL ^{7*}
Ketones	Negative	15 - 150 mg/dL (1+ - 3+)
Glucose	Normal	250 - 1000 mg/dL
Protein	Negative	30 - 100 mg/dL (1+ - 2+) ⁷
Nitrite	Negative	Positive
Leukocytes	Negative	100 - 500 Leu/ μ L (1+ - 2+)
pH	5 - 6.5	7 - 9
Specific Gravity	1.015 - 1.025	1.000 - 1.015
ROCHE Cobas 6500 (cobas u 601)		
Blood	Negative - Trace	150 - 250 Ery/ μ L
Leukocytes	Negative	100 - 500 Leu/ μ L
Nitrite	Negative	Positive
Ketones	Negative	15 - 150 mg/dL
Glucose	Normal	250 - 1000 mg/dL
Protein	Negative ⁶	30 - 100 mg/dL ⁷
Urobilinogen	Normal	1 - 8 mg/dL ^{7*}
Bilirubin	Negative	3 - 6 mg/dL*
pH	5 - 6	7 - 9
Specific Gravity	1.019 - 1.030	1.011 - 1.021
ROCHE cobas u 411		
Blood	Negative - Trace	50 - 250 Ery/ μ L (3+ - 5+)
Bilirubin	Negative	3 - 6 mg/dL (2+ - 3+)*
Urobilinogen	Negative - Normal	1 - 8 mg/dL (1+ - 3+) ^{7*}
Ketones	Negative	15 - 150 mg/dL (1+ - 4+)
Glucose	Negative - Normal	250 - 1000 mg/dL (3+ - 4+)
Protein	Negative ⁶	30 - 100 mg/dL (2+ - 3+) ⁷
Nitrite	Negative	Positive
Leukocytes	Negative	100 - 500 Leu/ μ L (2+ - 3+)
pH	5 - 6.5	7 - 9
Specific Gravity	1.015 - 1.025	1.000 - 1.020
ROCHE Urisys 1800		
Blood	Negative - Trace	25 - 250 Ery/ μ L (2+ - 5+)
Bilirubin	Negative	3 - 6 mg/dL (2+ - 3+)*
Urobilinogen	Negative - Normal	1 - 8 mg/dL (1+ - 3+) ^{7*}
Ketones	Negative	15 - 150 mg/dL (1+ - 4+)
Glucose	Negative - Normal	300 - 1000 mg/dL (3+ - 4+)
Protein	Negative ⁶	75 - 500 mg/dL (2+ - 4+) ⁷
Nitrite	Negative	Positive
Leukocytes	Negative	100 - 500 Leu/ μ L (2+ - 3+)
pH	5 - 6.5	7 - 9
Specific Gravity	1.015 - 1.025	1.000 - 1.020
ROCHE Urisys 2400		
Blood	Negative - Trace	25 - 250 Ery/ μ L
Bilirubin	Negative	3 - 6 mg/dL*
Urobilinogen	Normal	1 - 8 mg/dL ^{8*}
Ketones	Negative	15 - 150 mg/dL
Glucose	Normal	300 - 1000 mg/dL
Protein	Negative ⁶	25 - 150 mg/dL ⁸
Nitrite	Negative	Positive
Leukocytes	Negative	100 - 500 Leu/ μ L
pH	5 - 6.5	7 - 9
Specific Gravity	1.018 - 1.029	1.010 - 1.021

Analytes	Level 1 - 55251	Level 2 - 55252
SIEMENS VISUAL TESTING (Visual Test Strips Only)		
Glucose	Negative	100 - 1000 mg/dL
Bilirubin	Negative	Small - Large (1+ - 3+)
Ketones	Negative	Trace(5) - 160 mg/dL (Tr - Lg)
Specific Gravity	1.015 - 1.030	1.000 - 1.015
Blood	Negative - Trace	Trace - Large (Tr - 3+)
pH	5 - 6.5	7.0 - 8.5
Protein	Negative	30 - 300 mg/dL (1+ - 3+)
Urobilinogen	Normal (0.2 E.U./dL) ⁶	1.0 - 8.0 E.U./dL*
Nitrite	Negative	Positive
Leukocytes	Negative	Trace - Large (Tr - 3+)
Creatinine ²	10 - 100 mg/dL	100 - 300 mg/dL
SIEMENS® CLINITEK 50		
Glucose	Negative	100 - 500 mg/dL (Tr - 2+)
Bilirubin	Negative	Small - Large (1+ - 3+)
Ketones	Negative	Trace - \geq 80 mg/dL (\pm - 3+)
Specific Gravity	1.010 - \geq 1.030	\leq 1.005 - 1.020
Blood	Negative - Trace	Small - Large (1+ - 3+)
pH	5.0 - 6.5	7 - \geq 9
Protein	Negative	30 - \geq 300 mg/dL (1+ - 3+)
Urobilinogen	Normal (0.2 E.U./dL)	1.0 - 4.0 E.U./dL
Nitrite	Negative	Positive
Leukocytes	Negative	Trace - Large (Tr - 3+)
Microalbumin ¹	\leq 10 - 30 mg/L	30 - 300 mg/L
Creatinine ³	10 - 100 mg/dL	100 - 300 mg/dL
SIEMENS® CLINITEK 500		
Glucose	Negative	100 - 500 mg/dL (Tr - 2+)
Bilirubin	Negative	Small - Large (1+ - 3+)
Ketones	Negative	Trace - \geq 80 mg/dL (\pm - 3+)
Specific Gravity	1.015 - \geq 1.030	\leq 1.005 - 1.020
Blood	Negative - Trace	Small - Large (1+ - 3+)
pH	5 - 6.5	7.5 - \geq 9.0
Protein	Negative	30 - \geq 300 mg/dL (1+ - 3+)
Urobilinogen	Normal (0.2 E.U./dL)	1.0 - \geq 8.0 E.U./dL
Nitrite	Negative	Positive
Leukocytes	Negative	Trace - Large (Tr - 3+)
Creatinine ²	10 - 100 mg/dL	100 - 300 mg/dL
SIEMENS® CLINITEK ADVANTUS		
Glucose	Negative	100 - 500 mg/dL (Tr - 2+)
Bilirubin	Negative	Small - Large (1+ - 3+)
Ketones	Negative	Trace - \geq 80 mg/dL (Tr - 3+)
Specific Gravity	1.015 - \geq 1.030	\leq 1.005 - 1.020
Blood	Negative - Trace	Small - Large (1+ - 3+)
pH	5.0 - 6.5	7.0 - \geq 9.0
Protein	Negative	30 - \geq 300 mg/dL (1+ - 3+)
Urobilinogen	Normal (0.2 E.U./dL)	2.0 - \geq 8.0 E.U./dL
Nitrite	Negative	Positive
Leukocytes	Negative	Trace - Large (Tr - 3+)
Creatinine ²	10 - 100 mg/dL	100 - 300 mg/dL
SIEMENS CLINITEK STATUS/ STATUS PLUS/STATUS CONNECT		
Glucose	Negative	100 - 500 mg/dL (Tr - 2+)
Bilirubin	Negative	Small - Large (1+ - 3+)
Ketones	Negative	Trace - \geq 160 mg/dL (Tr - 4+)
Specific Gravity	1.010 - \geq 1.030	\leq 1.005 - 1.025
Blood	Negative - Trace	Small - Large (1+ - 3+)
pH	5.0 - 6.5	7 - \geq 9
Protein	Negative	30 - \geq 300 mg/dL (1+ - 3+)
Urobilinogen	Normal (0.2 E.U./dL)	2.0 - \geq 8.0 E.U./dL
Nitrite	Negative	Positive
Leukocytes	Negative	Trace - Large (Tr - 3+)
Microalbumin ¹	\leq 10 - 30 mg/L	30 - 300 mg/L
Creatinine ³	10 - 100 mg/dL	100 - 300 mg/dL
hCG	Negative	Positive
Teco Diagnostics TC-101 • TC-201 • TC-720 Urine Analyzer		

DATA NOT AVAILABLE AT THE TIME OF PRINTING.

Analytes	Level 1 - 55251	Level 2 - 55252
Uriscan™ • 10 SGL Strips (Visual)		
Blood	Negative - Trace	10 - 250 RBC/µL (1+ - 3+)
Bilirubin	Negative	0.5 - 3.0 mg/dL (1+ - 3+)
Urobilinogen	Negative - Normal	1 - 12 mg/dL (1+ - 4+)
Ketones	Negative	5 - 100 mg/dL (± - 3+)
Protein	Negative	30 - 300 mg/dL (1+ - 3+) ⁷
Nitrite	Negative	Positive
Glucose	Negative	100 - 1000 mg/dL (± - 3+)
pH	5.0 - 6.0	7.0 - 9.0
Specific Gravity	1.015 - 1.030	1.000 - 1.015
Leukocytes	Negative	25 - 500 WBC/µL (1+ - 3+)
Uriscan™ Optima Urine Analyzers • 10 SGL Strips		
Blood	Negative - Trace	10 - 250 RBC/µL (1+ - 3+)
Bilirubin	Negative	0.5 - 3.0 mg/dL (1+ - 3+)
Urobilinogen	Negative - Normal	1 - 12 mg/dL (1+ - 4+)
Ketones	Negative	5 - 100 mg/dL (± - 3+)
Protein	Negative	10 - 300 mg/dL (± - 3+) ⁷
Nitrite	Negative	Positive
Glucose	Negative	100 - 1000 mg/dL (± - 3+)
pH	5.0 - 6.5	6.5 - 8.0
Specific Gravity	1.015 - 1.030	1.005 - 1.020
Leukocytes	Negative	25 - 500 WBC/µL (1+ - 3+)

INTERNATIONAL USE ONLY

This Section is for International Use only and contains data for methods that are not available or cleared for diagnostic use in the United States.

Method	Level 1 - 55251	Level 2 - 55252	Units
Red Blood Cells (Erythrocytes)			
77 Elektronika UriSed Variants / COBIO Variants	0 - 30	0 - 150	p/µL
Beckman Coulter IRIS Diagnostics IQ@200 Series Analyzers	0 - 10	8 - 43	/µL
Mindray EH-2030, EH-2050-A Plus, EH-2050-B Plus, EH-2080B, EH-2080-C	0 - 20	0 - 60	Cell/µL
YD Diagnostics URISCAN PLUSCOPE	0 - 50	0 - 200	p/µL
ROCHE cobas 6500 (cobas u 701)	0 - 50	22 - 69	p/µL
KOVA® GLASSTIC® SLIDE 10 with GRIDS	3 - 13	18 - 95	/µL
Non-grid slides (~0.5 mL)	0 - 5	4 - 38	/hpf
Non-grid slides (~1.0 mL)	0 - 4	1 - 25	/hpf
Cen-Slide	0 - 5	3 - 40	/hpf
Slide & Coverslip (~0.5 mL)	0 - 6	3 - 21	/hpf
Slide & Coverslip (~1.0 mL)	0 - 4	1 - 13	/hpf
FisherBrand UriSystem DeciSlide	0 - 5	4 - 31	/hpf
White Blood Cells (Leukocytes)			
77 Elektronika UriSed Variants / COBIO Variants	0 - 30	20 - 90	p/µL
Beckman Coulter IRIS Diagnostics IQ@200 Series Analyzers	2 - 15	25 - 71	/µL
Mindray EH-2030, EH-2050-A Plus, EH-2050-B Plus, EH-2080B, EH-2080-C	0 - 20	0 - 100	Cell/µL
YD Diagnostics URISCAN PLUSCOPE	0 - 50	0 - 200	p/µL
ROCHE cobas 6500 (cobas u 701)	0 - 25	35 - 106	p/µL
KOVA® GLASSTIC® SLIDE 10 with GRIDS	2 - 15	17 - 85	/µL
Non-grid slides (~0.5 mL)	0 - 9	1 - 34	/hpf
Non-grid slides (~1.0 mL)	0 - 4	1 - 19	/hpf
Cen-Slide	0 - 6	0 - 41	/hpf
Slide & Coverslip (~0.5 mL)	0 - 4	1 - 19	/hpf
Slide & Coverslip (~1.0 mL)	0 - 4	0 - 10	/hpf
FisherBrand UriSystem DeciSlide	0 - 5	2 - 22	/hpf
Casts			
77 Elektronika UriSed Variants / COBIO Variants	None	None	
Mindray EH-2030, EH-2050-A Plus, EH-2050-B Plus, EH-2080B, EH-2080-C	None	None	
YD Diagnostics URISCAN PLUSCOPE	None	None	
KOVA® GLASSTIC® SLIDE 10 with GRIDS	None	None	
Non-grid slides (~0.5 mL)	None	None	
Non-grid slides (~1.0 mL)	None	None	
Cen-Slide	None	None	
Slide & Coverslip (~0.5 mL)	None	None	
Slide & Coverslip (~1.0 mL)	None	None	
FisherBrand UriSystem DeciSlide	None	None	

Method	Level 1 - 55251	Level 2 - 55252	Units
Crystals			
77 Elektronika UriSed Variants / COBIO Variants	may be present	present	
Beckman Coulter IRIS Diagnostics IQ@200 Series Analyzers	none	present	
Mindray EH-2030, EH-2050-A Plus, EH-2050-B Plus, EH-2080B, EH-2080-C	none	may be present	
YD Diagnostics URISCAN PLUSCOPE	none	may be present	
ROCHE cobas 6500 (cobas u 701)	negative	positive	
KOVA® GLASSTIC® SLIDE 10 with GRIDS	none	present	
Non-grid slides (~0.5 mL)	none	present	
Non-grid slides (~1.0 mL)	none	present	
Cen-Slide	none	present	
Slide & Coverslip (~0.5 mL)	none	present	
Slide & Coverslip (~1.0 mL)	none	present	
FisherBrand UriSystem DeciSlide	none	present	
Bacteria			
77 Elektronika UriSed Variants / COBIO Variants	may be present	present	
Beckman Coulter IRIS Diagnostics IQ@200 Series Analyzers	may be present	may be present	
KOVA® GLASSTIC® SLIDE 10 with GRIDS	may be present	present	
Non-grid slides (~0.5 mL)	may be present	present	
Non-grid slides (~1.0 mL)	may be present	present	
Cen-Slide	may be present	present	
Slide & Coverslip (~0.5 mL)	may be present	present	
Slide & Coverslip (~1.0 mL)	may be present	present	
FisherBrand UriSystem DeciSlide	may be present	present	

Analytes	Level 1 - 55251	Level 2 - 55252
77 Elektronika (Visual / Analyzers)		
Bilirubin	Negative	1 - 6 mg/dL (1+ - 3+) ⁸
Urobilinogen	Normal	2 - 12 mg/dL (1+ - 4+) ⁸
Ketones	Negative	5 - 150 mg/dL ((+) - 3+)
Ascorbic Acid	Negative	Negative
Glucose	Normal	50 - 1000 mg/dL (1+ - 4+)
Protein	Negative	30 - 500 mg/dL (1+ - 3+)
Blood	Negative-10 Ery/µL (Neg - 1+)	10 - 300 Ery/µL (1+ - 3+)
pH	5 - 6	7 - 8
Nitrite	Negative	Positive
Leukocytes	Negative	75 - 500 Leu/µL (2+ - 3+)
Specific Gravity	1.010 - 1.030	1.000 - 1.025
Analyticon® Combi Screen (Visual)		
Bilirubin	Negative	1+ - 3+
Urobilinogen	Normal	2 - 12 mg/dL (35 - 200 µmol/l) ^{8*}
Ketones	Negative	(+) - 3+
Ascorbic Acid	Negative	Negative
Glucose	Normal	50 - 1000 mg/dL (2.8 - 56 mmol/L)
Protein	Negative	30 - 500 mg/dL
Blood	Negative	10 - 300 Ery/µL (1+ - 3+)
pH	5 - 6	6 - 8
Nitrite	Negative ⁶	Positive
Leukocytes	Negative	25 - 500 Leu/µL
Specific Gravity	1.010 - 1.020	1.000 - 1.010
Creatinine	10 - 200 mg/dL (0.9 - 17.7 mmol/L)	200 - 300 mg/dL (17.7 - 26.5 mmol/L)
Microalbumin	10 - 80 mg/L	150 - 500 mg/dL
Analyticon® (Analyzers)		
Bilirubin	Negative	1 - 4 mg/dL 17-70 µmol/L (1+ - 3+)
Urobilinogen	Normal	2 - 12 mg/dL 35-200 µmol/L (1+ - 4+) ^{8*}
Ketones	Negative	10 - 300 mg/dL 1.0 - 30 mmol/L (+) - 3+
Ascorbic Acid	Negative - 20 mg/dL Negative - 1+	Negative - 20 mg/dL Negative - 1+
Glucose	Normal	50 - 1000 mg/dL 2.8 - 5.6 mmol/L (1+ - 5+)
Protein	Negative	30 - 500 mg/dL 0.3 - 5 g/L (1+ - 3+)
Blood	Negative	10 - 300 Ery/µL (1+ - 3+)
pH	5 - 7	6 - 9
Nitrite	Negative ⁶	Positive
Leukocytes	Negative	25 - 500 Leu/µL (1+ - 3+)
Specific Gravity	1.010 - 1.030	1.000 - 1.020

Analytes	Level 1 - 55251	Level 2 - 55252
CYPRESS DIAGNOSTICS Urine Strips • CYANStrip • CYANStrip Mini • Visual		
Urobilinogen	Normal (0.1-1mg/dL)	Normal-2 mg/dL (0.1-33 µmol/L) ¹¹
Glucose	Negative	50-2000 mg/dL (2.8-111 mmol/L)
Bilirubin	Negative	Small-Large (1+-3+)
Ketones	Negative	5-40 mg/dL (0.5-4 mmol/L) (Trace-2+)
Specific Gravity	1.015-1.025	1.005-1.020
Blood	Negative-Trace	10-250 RBC/µL (1+-3+)
pH	5-6.5	7-9
Protein	Negative	15-300 mg/dL (0.15-3.0 g/L) (Trace-3+)
Nitrite	Negative	Positive
Leukocytes	Negative	15-500 WBC/µL (Trace-3+)
Microalbumin	10-50 mg/dL (0.5-0.9 mmol/L)	100-300 mg/dL (8.8-26.5 mmol/L)
Creatinine	10 mg/L	30-150 mg/L
DiaLab Urine Strip Analyzer 500/Urine Strip 10C/Urine Strip 2MC		
Leukocytes	Negative	70-500 Leu/µL
Nitrite	Negative	Positive
Urobilinogen	0.2 - 1 mg/dL	0.2 - 4 mg/dL ¹¹
Protein	Negative	30 - 300 mg/dL
pH	5.0 - 7.0	7.0 - 9.0
Blood	Negative	1+ - 3+
Specific Gravity	1.015 - 1.030	1.005 - 1.025
Ketones	Negative	5 - 160 mg/dL
Bilirubin	Negative	1 - 4 mg/dL
Glucose	Negative	100 - 500 mg/dL
Creatinine	10 - 50 mg/dL	50 - 200 mg/dL
Microalbumin	1 - 3 mg/dL	3 - 15 mg/dL
DFI CYBOW • ComboStik • DUS Urine Reagent Strips (Visual)		
Urobilinogen	Normal (0.1-1 mg/dL)	Normal-2 mg/dL (0.1-33 µmol/L) ¹¹
Glucose	Negative	50-2000 mg/dL (2.8-111 mmol/L)
Bilirubin	Negative	Small-Large (1+-3+)
Ketones	Negative	5-40 mg/dL (0.5-4 mmol/L) (Trace-2+)
Specific Gravity	1.015-1.025	1.005-1.020
Blood	Negative-Trace	10-250 RBC/µL (1+-3+)
pH	5-6.5	7-9
Protein, Total	Negative	15-300 mg/dL (0.15-3.0 g/L) (Trace-3+)
Nitrite	Negative	Positive
Leukocytes	Negative	15-500 WBC/µL (Trace-3+)
Creatinine	10-50 mg/dL (0.5-0.9 mmol/L)	100-300 mg/dL (8.8-26.5 mmol/L)
Microalbumin	10 mg/L	30-150 mg/L
DFI CYBOW R-50 (50S) • ComboStik R-50 (50S) • DUS R-50 (50S)		
Urobilinogen	Normal (0.1-1 mg/dL)	Normal-2 mg/dL (0.1-33 µmol/L) ¹¹
Glucose	Negative	50-2000 mg/dL (2.8-111 mmol/L)
Bilirubin	Negative	Small-Large (1+-3+)
Ketones	Negative	5-40 mg/dL (0.5-4 mmol/L) (Trace-2+)
Specific Gravity	1.015-1.025	1.005-1.020
Blood	Negative-Trace	10-250 RBC/µL (1+-3+)
pH	5-6.5	7-9
Protein, Total	Negative	15-300 mg/dL (0.15-3.0 g/L) (Trace-3+)
Nitrite	Negative	Positive
Leukocytes	Negative	15-500 WBC/µL (Trace-3+)
Creatinine	10-50 mg/dL (0.5-0.9 mmol/L)	100-300 mg/dL (8.8-26.5 mmol/L)
Microalbumin	10 mg/L	30-150 mg/L
DFI CYBOW Reader 300 • ComboStik R-300 • DUS R-300		
Urobilinogen	Normal (0.1-1 mg/dL)	Normal-2 mg/dL (0.1-33 µmol/L) ¹¹
Glucose	Negative	50-2000 mg/dL (2.8-111 mmol/L)
Bilirubin	Negative	Small-Large (1+-3+)
Ketones	Negative	5-40 mg/dL (0.5-4 mmol/L) (Trace-2+)
Specific Gravity	1.015-1.025	1.005-1.020
Blood	Negative-Trace	10-250RBC/µL (1+-3+)
pH	5-6.5	7-9
Protein, Total	Negative	15-300mg/dL (0.15-3.0g/L) (Trace-3+)
Nitrite	Negative	Positive
Leukocytes	Negative	15-500 WBC/µL (Trace-3+)
Creatinine	10-50 mg/dL (0.5-0.9 mmol/L)	100-300 mg/dL (8.8-26.5 mmol/L)
Microalbumin	10 mg/L	30-150 mg/L

Analytes	Level 1 - 55251	Level 2 - 55252
DFI CYBOW Reader 720 • Combostik R-700 • DUS R-720 • DFI CYBOW Reader 720		
Urobilinogen	Normal (0.1-1 mg/dL)	Normal-2 mg/dL (0.1-33 µmol/L) ¹¹
Glucose	Negative	50-2000 mg/dL (2.8-111 mmol/L)
Bilirubin	Negative	Small-Large (1+-3+)
Ketones	Negative	5-40 mg/dL (0.5-4 mmol/L) (Trace-2+)
Specific Gravity	1.015-1.025	1.005-1.020
Blood	Negative-Trace	10-250RBC/µL (1+-3+)
pH	5-6.5	7-9
Protein, Total	Negative	15-300mg/dL (0.15-3.0g/L) (Trace-3+)
Nitrite	Negative	Positive
Leukocytes	Negative	15-500 WBC/µL (Trace-3+)
Creatinine	10-50 mg/dL (0.5-0.9 mmol/L)	100-300 mg/dL (8.8-26.5 mmol/L)
Microalbumin	10mg/L	30-150 mg/L
ERBA LACHEMA Dekaphan LAURA STRIPS & LAURA Urine Analyzer ERBA Mannheim Uro-dip 10e STRIPS & Uro-dipcheck 400e Urine Analyzer		
Bilirubin	Negative	3 - 6 mg/dL (51 - 103 µmol/L) (2+ - 3+)
Blood	Negative - Trace	50 - 250 Ery/µL (2+ - 3+)
Glucose	Negative	100 - 1000 mg/dL (5.5 - 55 mmol/L) (2+ - 4+)
Ketones	Negative	52 - 156 mg/dL (5 - 15 mmol/L) (2+ - 3+)
Leukocytes	Negative	75 - 500 Leu/µL (2+ - 3+)
Nitrite	Negative	Positive
pH	5 - 6	6.5 - 8
Protein	Negative	30 - 500 mg/dL (0.3 - 5 g/L) (1+ - 3+)
Specific Gravity	1,020 - 1,030	1.000 - 1.015
Urobilinogen	Normal	1 - 12 mg/dL (17 - 203 µmol/L) (1+ - 4+) ⁸
ERBA LACHEMA Dekaphan LAURA STRIPS & LAURA M Urine Analyzer ERBA Mannheim Uro-dip 10e STRIPS & LAURA M Urine Analyzer		
Bilirubin	Negative	3 - 6 mg/dL (51 - 103 µmol/L) (2+ - 3+)
Blood	Negative - Trace	50 - 250 Ery/µL (2+ - 3+)
Glucose	Negative	100 - 1000 mg/dL (5.5 - 55 mmol/L) (2+ - 4+)
Ketones	Negative	16 - 156 mg/dL (1.5 - 15 mmol/L) (1+ - 3+)
Leukocytes	Negative	75 - 500 Leu/µL (2+ - 3+)
Nitrite	Negative	Positive
pH	≤6	≤6 - 8
Protein	Negative	30 - 500 mg/dL (0.3 - 5 g/L) (1+ - 3+)
Specific Gravity	1,015 - 1,030	1.000 - 1.015
Urobilinogen	Normal	1 - 12 mg/dL (17 - 203 µmol/L) (1+ - 4+) ⁸
ERBA LACHEMA Dekaphan LAURA STRIPS & LAURA Smart Urine Analyzer ERBA Mannheim Uro-dip 10e STRIPS & Uro-dipcheck 240e Urine Analyzer		
Bilirubin	Negative	3 - 6 mg/dL (51 - 103 µmol/L) (2+ - 3+)
Blood	Negative - Trace	50 - 250 Ery/µL (2+ - 3+)
Glucose	Negative	100 - 1000 mg/dL (5.5 - 55 mmol/L) (2+ - 4+)
Ketones	Negative	52 - 156 mg/dL (5 - 15 mmol/L) (2+ - 3+)
Leukocytes	Negative	75 - 500 Leu/µL (2+ - 3+)
Nitrite	Negative	Positive
pH	5 - 6.5	6.5 - 8
Protein	Negative	30 - 500 mg/dL (0.3 - 5 g/L) (1+ - 3+)
Specific Gravity	1,020 - 1,030	1.000 - 1.015
Urobilinogen	Normal	1 - 12 mg/dL (17 - 203 µmol/L) (1+ - 4+) ⁸
ERBA LACHEMA Dekaphan LAURA STRIPS (Visual) ERBA Mannheim Uro-dip 10e STRIPS (Visual)		
Bilirubin	Negative	3 - 6 mg/dL (51 - 103 µmol/L) (2+ - 3+)
Blood	Negative - Trace	50 - 250 Ery/µL (2+ - 3+)
Glucose	Negative	300 - 1000 mg/dL (17 - 55 mmol/L) (3+ - 4+)
Ketones	Negative	52 - 156 mg/dL (5 - 15 mmol/L) (2+ - 3+)
Leukocytes	Negative	75 - 500 Leu/µL (2+ - 3+)
Nitrite	Negative	Positive
pH	5 - 6.5	6.5 - 8
Protein	Negative	30 - 500 mg/dL (0.3 - 5 g/L) (1+ - 3+)
Specific Gravity	1.025 - 1.030	1.000 - 1.015
Urobilinogen	Normal	1 - 12 mg/dL (17 - 203 µmol/L) (1+ - 4+) ⁸

Analytes	Level 1 - 55251	Level 2 - 55252
Mindray UA-5800/UA-6800		
Leukocytes	Negative	70-500 / μ L (1+~3+)
Urobilinogen	Normal	4-8 mg/dL (2+~3+)
Microalbumin	0.01-0.03 g/L	0.08-0.15 g/L
Protein	Negative	30-300 mg/dL (1+~3+)
Bilirubin	Negative	1-6 mg/dL (1+~3+)
Glucose	Negative	50-250 mg/dL (\pm 2+)
Specific Gravity	1.010-1.030	1.000-1.020
Ketones	Negative	15-80 mg/dL (1+~3+)
Nitrite	Negative	Positive
Creatinine	0.9-4.4 mmol/L	8.8-26.5 mmol/L
pH	5.0-6.5	6.5-8.0
Blood	Negative-Trace	25-200 / μ L (1+~3+)
ROCHE VISUAL TESTING (Visual Test Strips Only)		
Specific Gravity	1.015 - 1.030	1.000 - 1.010
pH	5 - 6	7 - 9
Leukocytes	Negative	75 - 500 Leu/ μ L (2+ - 3+)
Nitrite	Negative	Positive
Protein	Negative	30 - 100 mg/dL
Glucose	Normal	300 - 1000 mg/dL (3+ - 4+)
Ketones	Negative	10 - 150 mg/dL (1+ - 3+)
Urobilinogen ^{9*}	Normal	1 - 8 mg/dL (1+ - 3+)
Bilirubin ^{9*}	Negative	1+ - 3+
Blood	Negative - Trace	10 - 250 Ery/ μ L (1+ - 4+)
Microalbumin ⁶	Negative - 20 mg/L	50 - 100 mg/L
ROCHE cobas 6500 (cobas u 601)		
Blood	Negative - Trace	150 - 250 Ery/ μ L
Leukocytes	Negative	100 - 500 Leu/ μ L
Nitrite	Negative	Positive
Ketones	Negative	15 - 150 mg/dL
Glucose	Normal	300 - 1000 mg/dL
Protein	Negative ⁶	75 - 150 mg/dL ⁵
Urobilinogen	Normal	1 - 8 mg/dL ^{8*}
Bilirubin	Negative	3 - 6 mg/dL *
pH	5.0 - 6.0	7 - 9
Specific Gravity	1.019 - 1.030	1.011 - 1.022
ROCHE Miditron M		
Blood	Negative - Trace	150 - 250 Ery/ μ L
Bilirubin	Negative	3 - 6 mg/dL *
Urobilinogen	Normal	1 - 8 mg/dL ^{8*}
Ketones	Negative	15 - 150 mg/dL
Glucose	Normal	300 - 1000 mg/dL
Protein	Negative ⁶	25 - 150 mg/dL ⁵
Nitrite	Negative	Positive
Leukocytes	Negative	100 - 500 Leu/ μ L
pH	5 - 6.5	7 - 9
Specific Gravity	1.015 - 1.025	1.000 - 1.015
ROCHE Urisys 1100 Urine Analyzer or ROCHE Urilux S Urine Analyzer		
Blood	Negative - Trace	50 - 250 Ery/ μ L (3+ - 4+)
Bilirubin	Negative	3 - 6 mg/dL (2+ - 3+)*
Urobilinogen	Negative - Normal	1 - 8 mg/dL (1+ - 3+) ^{7*}
Ketones	Negative	5 - 150 mg/dL ((+) - 3+)
Glucose	Negative - Normal	300 - 1000 mg/dL (3+ - 4+)
Protein	Negative ⁵	25 - 150 mg/dL (1+ - 3+) ⁷
Nitrite	Negative	Positive (1+)
Leukocytes	Negative	100 - 500 Leu/ μ L (2+ - 3+)
pH	5 - 6.5	7 - 9
Specific Gravity	1.015 - 1.025	1.000 - 1.015
ROCHE Urisys 1800 Urine Analyzer or ROCHE Cobas u 411 Urine Analyzer		
Blood	Negative - Trace	150 - 250 Ery/ μ L (4+ - 5+)
Bilirubin	Negative	3 - 6 mg/dL (2+ - 3+)*
Urobilinogen	Negative - Normal	1 - 8 mg/dL (1+ - 3+) ^{8*}
Ketones	Negative	15 - 150 mg/dL (1+ - 4+)
Glucose	Negative - Normal	300 - 1000 mg/dL (3+ - 4+)
Protein	Negative ⁶	75 - 150 mg/dL (2+ - 3+) ⁵
Nitrite	Negative	Positive
Leukocytes	Negative	100 - 500 Leu/ μ L (2+ - 3+)
pH	5 - 6.5	7 - 9
Specific Gravity	1.015 - 1.025	1.000 - 1.020
ROCHE Urisys 2400		
Blood	Negative - Trace	150 - 250 Ery/ μ L
Bilirubin	Negative	3 - 6 mg/dL
Urobilinogen	Normal	1 - 8 mg/dL ^{8*}
Ketones	Negative	15 - 150 mg/dL
Glucose	Normal	300 - 1000 mg/dL
Protein	Negative ⁷	25 - 150 mg/dL ⁵
Nitrite	Negative	Positive
Leukocytes	Negative	100 - 500 Leu/ μ L
pH	5 - 6.5	7 - 9
Specific Gravity	1.018 - 1.029	1.010 - 1.021

Analytes	Level 1 - 55251	Level 2 - 55252
YD Diagnostics URISCAN Urine Test Strips (Visual)		
Blood	Negative	10-250 RBC/ μ L (1+~3+)
Bilirubin	Negative	0.5-3.0 mg/dL (1+~3+) ⁷
Urobilinogen	Normal	1-12 mg/dL (1+~4+)
Ketones	Negative	10-100 mg/dL (1+~3+)
Protein	Negative	30-1000 mg/dL (1+~4+)
Nitrite	Negative	Positive
Glucose	Negative	250-2000 mg/dL (1+~4+)
pH	5.0-6.5	7.0-9.0
Specific gravity	1.015-1.030	1.005-1.020
Leucocytes	Negative	25-500 WBC/ μ L (1+~3+)
YD Diagnostics URISCAN Pro II • URISCAN Optima II • URISCAN Pro • URISCAN Optima		
Blood	Negative	10-250 RBC/ μ L (1+~3+)
Bilirubin	Negative	0.5-3.0 mg/dL (1+~3+) ⁷
Urobilinogen	Normal	1-12 mg/dL (1+~4+)
Ketones	Negative	10-100 mg/dL (1+~3+)
Protein	Negative	30-1000 mg/dL (1+~4+)
Nitrite	Negative	Positive
Glucose	Negative	250-2000 mg/dL (1+~4+)
pH	5.0-6.5	6.5-8.5
Specific gravity	1.010-1.030	1.005-1.020
Leucocytes	Negative	25-500 WBC/ μ L (1+~3+)
Microalbumin	Negative-30mg/L	30-150mg/L
Creatinine	10-100mg/dL	100-300mg/dL
YD Diagnostics URISCAN Super YD Diagnostics URISCAN Super+		
Blood	Negative	10-250 RBC/ μ L (1+~3+)
Bilirubin	Negative	0.5-3.0 mg/dL (1+~3+) ⁷
Urobilinogen	Normal	1-12 mg/dL (1+~4+)
Ketones	Negative	10-100 mg/dL (1+~3+)
Protein	Negative	30-1000 mg/dL (1+~4+)
Nitrite	Negative	Positive
Glucose	Negative	250-2000 mg/dL (1+~4+)
pH	5.0-6.5	6.5-8.5
Specific gravity	1.020-1.030	1.012-1.022
Leucocytes	Negative	25-500 WBC/ μ L (1+~3+)
YD URISCAN Super +		
Blood	Negative	10-250 RBC/ μ L (1+~3+)
Bilirubin	Negative	0.5-3.0 mg/dL (1+~3+) ⁷
Urobilinogen	Normal	1-12 mg/dL (1+~4+)
Ketones	Negative	10-100 mg/dL (1+~3+)
Protein	Negative	30-1000 mg/dL (1+~4+)
Nitrite	Negative	Positive
Glucose	Negative	100-1000 mg/dL (\pm 3+)
pH	5.0-6.5	6.5-8.5
Specific gravity	1.020-1.030	1.012-1.022
Leucocytes	Negative	25-500 WBC/ μ L (1+~3+)

Footnotes for values, Fußnoten für werte, Apostilles pour des valeurs, Note a piè di pagina per i valori, Notas al pie de la página para los valores

ENGLISH

- Values only apply to Clinitek Microalbumin Reagent Strips when read on the Clinitek 50 and Status.
- Values only apply to Multistix Pro™ Reagent Strips
- Values only apply to Multistix Pro and Clinitek Microalbumin Reagent Strips when read on Clinitek Urine Analyzers
- VISUAL: Some customers may obtain false negatives.
- Values apply to Chemstrip® Micral Reagent Strips
- Some customers may obtain false positives.
- Some customers may obtain false negatives.
- Atypical color
- Values only apply to Siemens Clinitek 50, 500
- Absorbance at 620 nm
- The urobilinogen reaction produces an atypical color which may result in a normal (0.2 E.U./dL) reading. Should this occur, a visual observation of the intensification of the pad color indicates a positive response.
- See Limitations

DEUTSCH

- Werte gelten nur für Clinitek Mikroalbumin-Reagenzstreifen wenn diese auf Clinitek 50 und Status
- Werte gelten nur für Multistix Pro™ Reagenzstreifen
- Werte gelten nur für Multistix Pro und Clinitek Mikroalbumin-Reagenzstreifen, wenn diese auf Clinitek Urin-Analysatoren gelesen werden
- VISUAL: Manche Kunden erhalten möglicherweise falsch negative Ergebnisse.
- Werte gelten für Chemstrip® Micral Reagenzstreifen
- Manche Kunden erhalten möglicherweise falsch positive Ergebnisse.
- Manche Kunden erhalten möglicherweise falsch negative Ergebnisse.
- Atypische Farbe
- Werte gelten nur für Siemens Clinitek 50, 500
- Absorption bei 620 nm
- Die Urobilinogen-Reaktion erzeugt eine atypische Farbe, die zu einem normalen Messwert (0,2 EU/dl) führen kann. In diesem Fall kann eine positive Reaktion anhand der sichtbar veränderten Farbsättigung des Testfeldes festgestellt werden.
- Siehe Einschränkungen

FRANÇAIS

- Valeurs s'appliquent uniquement aux bandes de réactif Clinitek micro-albumine lues sur Clinitek 50 et Status
- Valeurs s'appliquent uniquement aux bandes de réactif Multistix Pro™
- Valeurs s'appliquent uniquement aux bandes de réactif Multistix Pro et Clinitek micro-albumine lues sur Clinitek Analyseurs d'urine
- VISUAL: Certains clients sont susceptibles d'obtenir des faux négatifs.
- Valeurs s'appliquent aux bandes de réactif Chemstrip® Micral
- Certains clients sont susceptibles d'obtenir des faux positifs.
- Certains clients sont susceptibles d'obtenir des faux négatifs.
- Couleur atypique
- Valeurs s'appliquent uniquement aux Siemens Clinitek 50, 500
- Absorbance à 620 nm
- La réaction de l'urobilinogène produit une couleur atypique pouvant donner lieu à une lecture normale (0,2 unité Ehrlich/dl). Si cela se produit, l'observation visuelle de l'intensification de la couleur de la zone de test indique une réponse positive.
- Voir Limitations

ITALIANO

- 1 I valori si riferiscono esclusivamente alle Strisce reagenti per microalbumina Clinitek lette su Clinitek 50 e Status
- 2 I valori si riferiscono esclusivamente alle Strisce reagenti Multistix Pro™
- 3 I valori si riferiscono esclusivamente alle Strisce reagenti per microalbumina Multistix Pro e Clinitek lette su Clinitek Analizzatori urine
- 4 VISUAL: Alcuni pazienti possono ottenere risultati falsi negativi.
- 5 I valori si riferiscono alle Strisce reagenti Micral Chemstrip®
- 6 Alcuni pazienti possono ottenere risultati falsi positivi.
- 7 Alcuni pazienti possono ottenere risultati falsi negativi.
- 8 Colore atipico
- 9 I valori si riferiscono esclusivamente alle Siemens Clinitek 50, 500
- 10 Assorbanza a 620 nm
- 11 La reazione dell'urobilinogeno produce un colore atipico che può determinare una lettura normale (0,2 U.E./dl). In questo caso, se si nota visivamente un'intensificazione del colore del cuscinetto, questo indica una reazione positiva.

• Vedere limiti

ESPAÑOL

- 1 Los valores son aplicables únicamente a las tiras reactivas Clinitek Microalbumin cuando se leen en equipos Clinitek 50 y Status
- 2 Los valores son aplicables únicamente a las tiras reactivas Multistix Pro™
- 3 Los valores son aplicables únicamente a las tiras reactivas Multistix Pro y Clinitek Microalbumin cuando se leen en equipos Clinitek Analizadores de orina
- 4 VISUAL: Algunos pacientes pueden obtener resultados negativos falsos.
- 5 Los valores son aplicables a las tiras reactivas Chemstrip® Micral
- 6 Algunos pacientes pueden obtener resultados positivos falsos
- 7 Algunos pacientes pueden obtener resultados negativos falsos.
- 8 Color anormal
- 9 Los valores son aplicables únicamente a las Siemens Clinitek 50, 500
- 10 Absorbencia a 620 nm
- 11 La reacción del urobilinógeno genera un color atípico que puede dar lugar a una lectura normal (0,2 E.U./dl). Si ocurriera esto, la respuesta es positiva si se observa visualmente una intensificación del color de la almohadilla.

• Ver las limitaciones

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