

Adulterants Survey, Chain of Custody, Random, Urine

#### Overview

#### **Useful For**

Assess the possible adulteration of a urine specimen submitted for drug of abuse testing, as well as for providing the urine creatinine for "creatinine normalization"

Chain of custody is required whenever the results of testing could be used in a court of law. Its purpose is to protect the rights of the individual contributing the specimen by demonstrating that it was under the control of personnel involved with testing the specimen at all times; this control implies that the opportunity for specimen tampering would be limited.

#### **Additional Tests**

Test Id	Reporting Name	Available Separately	Always Performed
COCH	Chain of Custody	No	Yes
	Processing		

#### **Testing Algorithm**

See <u>Adulterant Survey Algorithm</u> in Special Instructions.

#### **Special Instructions**

Adulterant Survey Algorithm

#### **Method Name**

Spectrophotometry

## **NY State Available**

Yes

## Specimen

#### Specimen Type

Urine

## **Specimen Required**

**Supplies:** Chain of Custody Kit (T282)

**Container/Tube:** Chain-of-custody kit containing the specimen containers, seals, and documentation required.

Specimen Volume: 1.5 mL

Collection Instructions: Collect specimen in the provided container, seal, and submit with the associated documentation



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to satisfy the legal requirements for chain-of-custody testing.

Additional Information: Submitting less than 1.5 mL will compromise the ability to perform all necessary testing.

#### **Forms**

- 1. Chain of Custody Request is included in the Chain-of-Custody Kit (T282).
- 2. If not ordering electronically, complete, print, and send a Therapeutics Test Request (T831) with the specimen.

## **Specimen Minimum Volume**

1 mL

## Reject Due To

All specimens will be evaluated at Mayo Clinic Laboratories for test suitability.

## **Specimen Stability Information**

Specimen Type	Temperature	Time	Special Container
Urine	Refrigerated (preferred)	14 days	
	Frozen	14 days	
	Ambient	72 hours	

## **Clinical & Interpretive**

## **Clinical Information**

Specimen adulteration is the manipulation of a sample that may cause false-negative test results for the presence of drugs of abuse. Common adulterants that may affect testing are water, soap, bleach, vinegar, oxidants, and salt. The adulteration testing includes assessment of creatinine concentration, pH, urine specific gravity, presence or absence of an oxidant, and presence or absence of nitrite.

Chain of custody is a record of the disposition of a specimen to document each individual who collected the specimen, handled it, and performed the analysis. When a specimen is submitted in this manner, analysis will be performed in such a way that it will withstand regular court scrutiny.

## **Reference Values**

Cutoff concentrations Oxidants: 200 mg/L Nitrites: 500 mg/L

#### Interpretation

See Adulterant Survey Algorithm in Special Instructions.

#### **Cautions**

No significant cautionary statements



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#### Clinical Reference

US Department of Health and Human Services, Substance Abuse and Mental Health Services Administration (SAMHSA): Mandatory Guidelines for Federal Workplace Drug Testing Programs. Federal Register. 2017 January 23;82(13):FR 7920. Available at: www.samhsa.gov/sites/default/files/workplace/frn\_vol\_82\_7920\_.pdf
US Department of Health and Human Services, Substance Abuse and Mental Health Services Administration (SAMHSA): Drug-Free Workplace Guidelines and Resources: Substance Abuse and Mental Health Services Administration (SAMHSA). Updated June 7, 2021. Accessed September 27, 2021. Available at: www.samhsa.gov/workplace/resources.

#### **Performance**

#### **Method Description**

All results are measured using spectrophotometry at wavelengths specified by the reagent manufacturer. The use of a refractometer may also be used in the specific gravity measurement. (Package insert: Creatinine plus ver 2 Specimen Validity Test Nitrite, Specimen Validity Test Oxidant, Specimen Validity Test pH, Specimen Validity Test Specific Gravity, Roche Diagnostics; 12/2016)

## **PDF Report**

No

## Day(s) Performed

Monday through Saturday

#### Report Available

2 to 3 days

#### **Specimen Retention Time**

2 weeks

## **Performing Laboratory Location**

Rochester

#### Fees & Codes

#### Fees

- Authorized users can sign in to <u>Test Prices</u> for detailed fee information.
- Clients without access to Test Prices can contact <u>Customer Service</u> 24 hours a day, seven days a week.
- Prospective clients should contact their account representative. For assistance, contact <u>Customer Service</u>.

#### **Test Classification**

This test has been cleared, approved, or is exempt by the US Food and Drug Administration and is used per manufacturer's instructions. Performance characteristics were verified by Mayo Clinic in a manner consistent with CLIA



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requirements.

## **CPT Code Information**

81005

## **LOINC®** Information

Test ID	Test Order Name	Order LOINC® Value
ADLTX	Adulterants Survey, CoC, U	58715-4

Result ID	Test Result Name	Result LOINC® Value
36121	Creatinine, U	2161-8
36122	Specific Gravity	5810-7
36123	рН	2756-5
36124	Oxidants	58714-7
36125	Nitrites	32710-6
36126	Comment	48767-8