

1A  
1B  
1C  
Item# 0050659 Form # 260N.30

1A  
1B  
1C  
1D

**PHYSICIAN**

Physician/Authorized Signature: \_\_\_\_\_

Copy To: \_\_\_\_\_

Name \_\_\_\_\_

Address \_\_\_\_\_

City \_\_\_\_\_ State \_\_\_\_\_ Zip \_\_\_\_\_

Requesting Physician & NPI \_\_\_\_\_

**PATIENT**

Name (Last, First) \_\_\_\_\_ MI \_\_\_\_\_ MRN \_\_\_\_\_ DOB \_\_\_\_\_

Address \_\_\_\_\_ City \_\_\_\_\_ State \_\_\_\_\_ Zip \_\_\_\_\_

Home # ( ) \_\_\_\_\_ Work # ( ) \_\_\_\_\_  Male  Female Race:  Black  White  Hispanic  Other: \_\_\_\_\_

**BILLING**

Bill:  Medicare  Medicaid  Insurance  Patient  Ordering Physician  Facility (Account) Authorization # \_\_\_\_\_

Policy/ID # \_\_\_\_\_ Group # \_\_\_\_\_ 2nd Insurance Policy/ID # \_\_\_\_\_ Group # \_\_\_\_\_

Insurance Carrier \_\_\_\_\_ Insured's Name \_\_\_\_\_  
(if not patient check one -  spouse  child  other)

Claim Address \_\_\_\_\_ 2nd Insurance Carrier \_\_\_\_\_

City \_\_\_\_\_ State \_\_\_\_\_ Zip \_\_\_\_\_ Phone # \_\_\_\_\_ Claim Address \_\_\_\_\_

Patient Status  Hospital Inpatient  Hospital Outpatient  Hospital Non-Patient Insured's DOB \_\_\_\_\_

Billing Information Attached

**HISTOLOGY = Gross & Microscopic Exam**

**ICD-CM<sup>+</sup>**

Collection Date \_\_\_\_\_ Collection Time \_\_\_\_\_

Specimen Type \_\_\_\_\_

Number of Vials Submitted \_\_\_\_\_  
(UroScore<sup>®</sup> requires a sextant (6+ vials) biopsy & a PSA Value)

Prostate Histology  Prostate Histology w/UroScore<sup>®</sup>

Prostate Histology, Reflex to ProMark<sup>®</sup> Prognostic Test@ on:  
 Gleason 6  Gleason 7 (3+4)  Gleason 6 or 7 (3+4)  
ProMark<sup>®</sup> only available to CTR Certified physicians.

Prostate Histology, if Gleason 6 or 7 (3+4), Reflex to:  
 PTEN IHC  PTEN/ERG IHC

Prostate Histology, Reflex to ConfirmMDx@ on negative/HGPIN

Bladder Histology Biopsy

Bladder Histology TUR

Vas Deferens (Sterilization) Histology

Other Histology: \_\_\_\_\_

Consultation: \_\_\_\_\_

PSA Date \_\_\_\_\_ PSA \_\_\_\_\_ ng/mL

**DRE Finding**  Suspicious  Normal  
 Isolated Nodule  Multiple Nodules

**Previous Biopsy**  Positive  Negative  
 PIN  Suspicious

**Therapy**  Cryosurgery  Chemotherapy  
 Hormone Therapy  Radiation Therapy

**MicrocystePLUS<sup>®</sup> URINE CYTOLOGY**

**ICD-CM<sup>+</sup>**

Collection Date \_\_\_\_\_ Collection Time \_\_\_\_\_ AM   
PM

**Clinical Data**  Hematuria  
 TCC, Current  TCC, History Dx Date: \_\_\_\_\_  
 Other: \_\_\_\_\_

**Specimen Type (Required)**  
 Voided Urine (Bladder)  Catheterized Urine  
 Post-Cysto Void  Bladder Wash  
 Ileal Conduit/NeoBladder  Urethral Wash  
 Renal Wash - Left  Renal Wash - Right  
 Ureter Wash - Left  Ureter Wash - Right  
 Other: \_\_\_\_\_

**INDIVIDUAL TESTS: (May be ordered or added to profile)**  
 VU6 Pap Stain (only) Urine Cytology †  
 FNA (Fine Needle Aspiration) Site: \_\_\_\_\_  
 K600D Bladder Cancer FISH (Pathologist Review) †  
 β2 Microglobulin ✦  Microalbumin ✦  Total Protein ✦

**MicrocystePLUS<sup>®</sup> URINE CYTOLOGY PROFILES**

**994 Hematuria Profile** ✦  
Cytodiagnostic Urinalysis Correlating Cytology (by concentration technique, with Pap and Feulgen stain), Urine Dipstick, Microalbumin, β2 Microglobulin, Total Protein. 994 only on void, catheterized, or post-cysto void; other collection methods processed as U03 cytology.

**VU3 Cytology Plus Monitoring Profile** †  
Cytology (Pap and Feulgen stain)

**VU1D Bladder Cancer FISH/Cytology Pathodiagnostic Profile** †  
Bladder Cancer FISH Assay and Cytology (Pap and Feulgen Stain) Including integrated cytomolecular diagnostic interpretation with clinical correlation by pathologist (MD)

**VU4D Bladder Cancer FISH Reflex/Cytology Pathodiagnostic Profile** †  
Cytology (Pap and Feulgen stain); reflex to Bladder Cancer FISH (Pathologist review) on atypical cytology results

† TCC Monitoring Kit (Alcohol Fixative)  
✦ Urine Cytopathology Kit (Tablet Preservative)  
See reverse for collection method requirements and CPT codes

**CHEMISTRY**

**ICD-CM<sup>+</sup>**

Specimen Type \_\_\_\_\_

Collection Date \_\_\_\_\_ Collection Time \_\_\_\_\_ AM   
PM

**Fasting?** Yes  No  **Frozen**

S = Serum U = 24 Hr. Urine

**Endocrinology**

Total PSA@%  Alkaline Phosphatase  
 Total PSA@%/Rflx Free PSA  Albumin (S)  
with free/total PSA ratio  ALT  
 Total PSA@% and Free PSA  Ammonia (U)  
with free/total PSA ratio  AST  
 Testosterone  BUN  
 Unbound Testosterone  Calcium (S, U)  
 Testosterone/Unbound  Chloride (S, U)  
Testosterone with % Free  Cholesterol@%  
 FSH  Citrate (U)  
 LH  CO2  
 Prolactin  Creatinine (S, U)  
 AFP@  Cystine (U)\*  
 Beta HCG@%  Direct Bilirubin  
 TSH@%  Glucose@%  
 HDL@%  
 Magnesium (S, U)  
 Oxalate (U)  
 pH (U)

**Panels**

Electrolyte Panel  Phosphorus (S, U)  
 Lipid Panel@%  Potassium (S, U)  
 Hepatic Function Panel  PTH  
 Basic Metabolic Panel  Sodium (S, U)  
 Renal Function Panel  Sulfate (U)  
 Comp. Metabolic Panel  Total Bilirubin  
 Panel components on back  Total Protein (S, U)  
 Triglyceride@%  
 Uric Acid (S, U)

Other: \_\_\_\_\_

Dianon performed venipuncture & PST Initials \_\_\_\_\_

**24 HR URINE**

**ICD-CM<sup>+</sup>**

**REQUIRED**

Total Volume	Specimen Type	Collection Date	Collection Time
mils	Type		AM <input type="checkbox"/> PM <input type="checkbox"/>

**24 Hr Urine Chemistry Profiles (Dianon 24hr Urine Kit REQUIRED)**  
Check profile below or individual tests available in Chemistry section.

UroStone<sup>®</sup>24 Uric Acid (Uric Acid/Creatinine/Sulfate)

Creatinine Clearance (Serum Creatinine/Urine Creatinine)  
*requires serum & urine specimens and*

Patient Height: \_\_\_\_\_ Inches & Weight: \_\_\_\_\_ lbs.

UroStone<sup>®</sup>24 Cystine\* (Creatinine/Qualitative Cystine\*)

UroStone<sup>®</sup>24 Calcium (Creatinine/Calcium/Sodium/pH)

UroStone<sup>®</sup>24 Citrate (Citrate/Creatinine)

UroStone<sup>®</sup>24\* (Calcium/Citrate/Creatinine/Magnesium/  
Oxalate/pH/Phosphorus/Qualitative Cystine\*/Sodium/Uric Acid)

UroStone<sup>®</sup>Max24\* (Ammonia/Calcium/Chloride/Citrate/Creatinine/  
Magnesium/Oxalate/pH/Phosphorus/Potassium/Sodium/Sulfate  
Uric Acid/Qualitative Cystine\*)

**STONES**

**ICD-CM<sup>+</sup>**

Specimen Type \_\_\_\_\_

Collection Date \_\_\_\_\_ Collection Time \_\_\_\_\_ AM   
PM

**Stone Analysis - Urinary Tract Calculus** (Stone Analysis Kit)  
 Spontaneously Passed  
 Lithotripsy  
 Surgically Removed

**HISTORY**

Indicate previous Urinary Tract/Systemic Disorders, Biopsy or Therapy Results, and current Medications: \_\_\_\_\_

\* Quantitative Cystine performed on positive Qualitative Cystine at additional charge.  
When ordering tests for which Medicare or Medicaid reimbursement will be sought, physicians should only order tests that are medically necessary for the diagnosis or treatment of the patient.  
† Separately billable stains may be added by pathologist when medically necessary to render a diagnosis.  
(260N) Rev 12/16/2020  
Dianon Pathology is a brand of Dianon Systems, Inc., a wholly-owned subsidiary of Laboratory Corporation of America<sup>®</sup> Holdings.

Refer to Determining Necessity of ABN Completion on reverse.

**Symbols Legend**  
@ = Subject to Medicare medical necessity guidelines.  
% = Subject to Medicare frequency guidelines.  
# = Medicare deems investigational. Medicare does not pay for services it deems investigational.

**PLEASE ENSURE REQUESTING PHYSICIAN IS INDICATED AND THE TEST REQUESTED IS MARKED.**

**Specimen Label Instructions . . .**

1. Complete the requisition with all requested information.
2. Remove the required number of labels from the front of this sheet.
3. Place one (1) label on each specimen container (not on lid).

**Any Questions?**  
**Please Call Client Services at 1-800-411-1839**

SITE, IF APPLICABLE  
Dianon Pathology

SITE, IF APPLICABLE  
Dianon Pathology

**L TRANSITION ZONE**  
Dianon Pathology

**R TRANSITION ZONE**  
Dianon Pathology

**LEFT**  
Dianon Pathology

**RIGHT**  
Dianon Pathology

**LLAT BASE**  
Dianon Pathology

**L LAT MID**  
Dianon Pathology

**LLAT APEX**  
Dianon Pathology

**R LAT BASE**  
Dianon Pathology

**R LAT MID**  
Dianon Pathology

**R LAT APEX**  
Dianon Pathology

**LEFT BASE**  
Dianon Pathology

**LEFT MID**  
Dianon Pathology

**LEFT APEX**  
Dianon Pathology

**RIGHT BASE**  
Dianon Pathology

**RIGHT MID**  
Dianon Pathology

**RIGHT APEX**  
Dianon Pathology

2A  
2B  
2C  
Item# 0050659 Form # 260N.30

2A  
2B  
2C

PHYSICIAN  
Physician/Authorized Signature: \_\_\_\_\_  
Copy To: \_\_\_\_\_  
Name \_\_\_\_\_  
Address \_\_\_\_\_  
City \_\_\_\_\_ State \_\_\_\_\_ Zip \_\_\_\_\_  
Requesting Physician & NPI \_\_\_\_\_

PATIENT  
Name (Last, First) \_\_\_\_\_ MI \_\_\_\_\_ MRN \_\_\_\_\_ DOB \_\_\_\_\_  
Address \_\_\_\_\_ City \_\_\_\_\_ State \_\_\_\_\_ Zip \_\_\_\_\_  
Home # ( \_\_\_\_\_ ) \_\_\_\_\_ Work # ( \_\_\_\_\_ ) \_\_\_\_\_  
 Male  Female Race:  Black  White  Hispanic  Other: \_\_\_\_\_

BILLING  
Bill:  Medicare  Medicaid  Insurance  Patient  Ordering Physician  Facility (Account) Authorization # \_\_\_\_\_  
Policy/ID # \_\_\_\_\_ Group # \_\_\_\_\_ 2nd Insurance Policy/ID # \_\_\_\_\_ Group # \_\_\_\_\_  
Insurance Carrier \_\_\_\_\_ Attach secondary billing info. Insured's Name \_\_\_\_\_  
Claim Address \_\_\_\_\_ (if not patient check one -  spouse  child  other) 2nd Insurance Carrier \_\_\_\_\_  
City \_\_\_\_\_ State \_\_\_\_\_ Zip \_\_\_\_\_ Phone # \_\_\_\_\_ Claim Address \_\_\_\_\_  
Patient Status  Hospital Inpatient  Hospital Outpatient  Hospital Non-Patient Insured's DOB \_\_\_\_\_  
 Billing Information Attached

HISTOLOGY = Gross & Microscopic Exam  
ICD-CM<sup>▲</sup> \_\_\_\_\_  
Collection Date \_\_\_\_\_ Collection Time \_\_\_\_\_ AM  PM   
Specimen Type \_\_\_\_\_  
Number of Vials Submitted \_\_\_\_\_  
(UroScore<sup>®</sup> requires a sextant (6+ vials) biopsy & a PSA Value)  
 Prostate Histology  Prostate Histology w/UroScore<sup>®</sup>  
Prostate Histology, Reflex to ProMark<sup>®</sup> Prognostic Test@ on:  
 Gleason 6  Gleason 7 (3+4)  Gleason 6 or 7 (3+4)  
ProMark<sup>®</sup> only available to CTR Certified physicians.  
Prostate Histology, if Gleason 6 or 7 (3+4), Reflex to:  
 PTEN IHC  PTEN/ERG IHC  
 Prostate Histology, Reflex to ConfirmMDx@ on negative/HGPN  
 Bladder Histology Biopsy  
 Bladder Histology TUR  
 Vas Deferens (Sterilization) Histology  
 Other Histology: \_\_\_\_\_  
 Consultation: \_\_\_\_\_  
PSA Date \_\_\_\_\_ PSA \_\_\_\_\_ ng/mL  
DRE Finding  Suspicious  Normal  
 Isolated Nodule  Multiple Nodules  
Previous Biopsy  Positive  Negative  
 PIN  Suspicious  
Therapy  Cryosurgery  Chemotherapy  
 Hormone Therapy  Radiation Therapy

HISTOLOGY = Gross & Microscopic Exam  
ICD-CM<sup>▲</sup> \_\_\_\_\_  
Collection Date \_\_\_\_\_ Collection Time \_\_\_\_\_ AM  PM   
Clinical Data  Hematuria  
 TCC, Current  TCC, History Dx Date: \_\_\_\_\_  
 Other: \_\_\_\_\_  
Specimen Type (Required)  
 Voided Urine (Bladder)  Catheterized Urine  
 Post-Cysto Void  Bladder Wash  
 Ileal Conduit/NeoBladder  Urethral Wash  
 Renal Wash - Left  Renal Wash - Right  
 Ureter Wash - Left  Ureter Wash - Right  
 Other: \_\_\_\_\_  
INDIVIDUAL TESTS: (May be ordered or added to profile)  
 VU6 Pap Stain (only) Urine Cytology ‡  
 FNA (Fine Needle Aspiration) Site: \_\_\_\_\_  
 K600D Bladder Cancer FISH (Pathologist Review) ‡  
 β2 Microglobulin ✦  Microalbumin ✦  Total Protein ✦  
MicrocytePLUS<sup>®</sup> URINE CYTOLOGY PROFILES  
 994 Hematuria Profile ✦  
Cyodiagnostic Urinalysis Correlating Cytology (by concentration technique, with Pap and Feulgen stain), Urine Dipstick, Microalbumin, β2 Microglobulin, Total Protein. 994 only on void, catheterized, or post-cysto void; other collection methods processed as U03 cytology.  
 VU3 Cytology Plus Monitoring Profile ‡  
Cytology (Pap and Feulgen stain)  
 VU1D Bladder Cancer FISH/Cytology Pathodiagnostic Profile ‡  
Bladder Cancer FISH Assay and Cytology (Pap and Feulgen Stain) including integrated cytomolecular diagnostic interpretation with clinical correlation by pathologist (MD)  
 VU4D Bladder Cancer FISH Reflex/Cytology Pathodiagnostic Profile ‡  
Cytology (Pap and Feulgen stain); reflex to Bladder Cancer FISH (Pathologist review) on atypical cytology results  
‡ TCC Monitoring Kit (Alcohol Fixative)  
✦ Urine Cytopathology Kit (Tablet Preservative)  
See reverse for collection method requirements and CPT codes

CHEMISTRY  
ICD-CM<sup>▲</sup> \_\_\_\_\_  
Specimen Type \_\_\_\_\_  
Collection Date \_\_\_\_\_ Collection Time \_\_\_\_\_ AM  PM   
Fasting? Yes  No  Frozen   
S = Serum U = 24 Hr. Urine  
Endocrinology Individual Serum/  
24 Hr. Urine Chemistry  
 Total PSA@%  Alkaline Phosphatase  
 Total PSA@%/Rflx Free PSA  Albumin (S)  
with free/total PSA ratio  ALT  
 Total PSA@% and Free PSA  Ammonia (U)  
with free/total PSA ratio  AST  
 Testosterone  BUN  
 Unbound Testosterone  Calcium (S, U)  
 Testosterone/Unbound  Chloride (S, U)  
Testosterone with % Free  Cholesterol@%  
 FSH  Citrate (U)  
 LH  CO2  
 Prolactin  Creatinine (S, U)  
 AFP@  Cystine (U)★  
 Beta HCG@%  Direct Bilirubin  
 TSH@%  Glucose@%  
 HDL@%  
 Magnesium (S, U)  
 Oxalate (U)  
 pH (U)  
Panels  Electrolyte Panel  Phosphorus (S, U)  
 Lipid Panel@%  Potassium (S, U)  
 Hepatic Function Panel  PTH  
 Basic Metabolic Panel  Sodium (S, U)  
 Renal Function Panel  Sulfate (U)  
 Comp. Metabolic Panel  Total Bilirubin  
Panel components on back  Total Protein (S, U)  
 Triglyceride@%  
 Uric Acid (S, U)  
 Other: \_\_\_\_\_  
 Dianon performed venipuncture & PST Initials \_\_\_\_\_

24 HR URINE  
ICD-CM<sup>▲</sup> \_\_\_\_\_  
REQUIRED  
Total Volume \_\_\_\_\_ mls Specimen Type \_\_\_\_\_  
Collection Date \_\_\_\_\_ Collection Time \_\_\_\_\_ AM  PM   
24 Hr Urine Chemistry Profiles (Dianon 24hr Urine Kit REQUIRED)  
Check profile below or individual tests available in Chemistry section.  
 UroStone<sup>®</sup>24 Uric Acid (Uric Acid/Creatinine/Sulfate)  
 Creatinine Clearance (Serum Creatinine/Urine Creatinine)  
*requires serum & urine specimens and*  
Patient Height: \_\_\_\_\_ Inches & Weight: \_\_\_\_\_ lbs.  
 UroStone<sup>®</sup>24 Cystine★ (Creatinine/Qualitative Cystine\*)  
 UroStone<sup>®</sup>24 Calcium (Creatinine/Calcium/Sodium/pH)  
 UroStone<sup>®</sup>24 Citrate (Citrate/Creatinine)  
 UroStone<sup>®</sup>24★ (Calcium/Citrate/Creatinine/Magnesium/  
Oxalate/pH/Phosphorus/Qualitative Cystine\*/Sodium/Uric Acid)  
 UroStone<sup>®</sup>Max24★ (Ammonia/Calcium/Chloride/Citrate/Creatinine/  
Magnesium/Oxalate/pH/Phosphorus/Potassium/Sodium/Sulfate  
Uric Acid/Qualitative Cystine\*)

STONES  
ICD-CM<sup>▲</sup> \_\_\_\_\_  
Specimen Type \_\_\_\_\_  
Collection Date \_\_\_\_\_ Collection Time \_\_\_\_\_ AM  PM   
 Stone Analysis - Urinary Tract Calculus (Stone Analysis Kit)  
 Spontaneously Passed  
 Lithotripsy  
 Surgically Removed

HISTORY  
Indicate previous Urinary Tract/Systemic Disorders, Biopsy or Therapy Results, and current Medications: \_\_\_\_\_

\* Quantitative Cystine performed on positive Qualitative Cystine at additional charge.  
When ordering tests for which Medicare or Medicaid reimbursement will be sought, physicians should only order tests that are medically necessary for the diagnosis or treatment of the patient. PHYSICIAN'S COPY (260N) Rev 12/16/2020  
† Separately billable stains may be added by pathologist when medically necessary to render a diagnosis. Dianon Pathology is a brand of Dianon Systems, Inc., a wholly-owned subsidiary of Laboratory Corporation of America® Holdings.

Refer to Determining Necessity of ABN Completion on reverse.  
Symbols Legend  
@ = Subject to Medicare medical necessity guidelines.  
% = Subject to Medicare frequency guidelines.  
# = Medicare deems investigational. Medicare does not pay for services it deems investigational.

PLEASE ENSURE REQUESTING PHYSICIAN IS INDICATED AND THE TEST REQUESTED IS MARKED.

Specimen Label Instructions . . .

- 1. Complete the requisition with all requested information.
- 2. Remove the required number of labels from the front of this sheet.
- 3. Place one (1) label on each specimen container (not on lid).

Any Questions?  
Please Call Client Services  
at 1-800-411-1839



**Medical Necessity**

**Determining Necessity of Advance Beneficiary Notice of Non-coverage (ABN) Completion\***

- 1. Diagnose.** Determine your patient's diagnosis.
  - 2. Document.** Write the diagnosis code(s) on the front of this requisition.
  - 3. Verify.** Determine if the laboratory test(s) ordered for the patient is subject to Local Coverage Determination or National Coverage Determination. This information can be located in the policies published by your Medicare Administrative Contractor (MAC), CMS, or [www.Labcorp.com/MedicareMedicalNecessity](http://www.Labcorp.com/MedicareMedicalNecessity).
  - 4. Review.** If the diagnosis code for your patient does not meet the medical necessity requirements set forth by the Medicare carrier or the test(s) is/are being performed more frequently than the carrier allows, an ABN should be completed.
- \*An ABN should be completed for all tests that are considered research or investigational by Medicare.

**How to Complete an Advance Beneficiary Notice of Non-coverage (ABN)**

Medicare is very specific in requiring that all of the information included on the ABN be completed. Additionally, Labcorp requests that the specimen number or bar code label be included on the form. To be valid an ABN must:

1. Be executed on the CMS approved ABN form (CMS-R-131)
2. Identify the Medicare Part B Beneficiary, using the name as it appears on the patient's red, white and blue Medicare card
3. Indicate the test(s)/procedure(s) which may be denied within the relevant reason column
4. Include an estimated cost for the test(s)/procedure(s) subject to the ABN
5. Have 'Option 1', 'Option 2', or 'Option 3' designated by the beneficiary
6. Be signed **and** dated by the beneficiary or his/her representative **prior to** the service being rendered

**Symbols used to designate local/national medical review as of 10/01/2020**

- @ = Subject to Medicare medical necessity guidelines
- % = Subject to Medicare frequency guidelines
- # = Medicare deems investigational. Medicare does not pay for services it deems investigational.

**TUBE AND SPECIMEN TRANSPORTATION REQUIREMENTS**

TEST	TUBE	CPT	SPECIMEN	TEST	TUBE	CPT	SPECIMEN	TEST	TUBE	CPT	SPECIMEN
AFP	(SST)	82105	(S,R)	Comprehensive Metabolic Panel	(SST)	80053	(S,R)	Prolactin	(SST)	84146	(S,R)
Albumin	(SST)	82040	(S,R)	Creatinine	(SST)	82565	(S,R)	PSA	(SST)	84153	(S,R)
ALT	(SST)	84460	(S,R)	Creatinine Clearance	(Urine+SST)	82575	(U,S,R)	PSA, Free	(SST)	84154	(S,R)
Alkaline Phosphatase	(SST)	84075	(S,R)	Direct Bilirubin	(SST)	82248	(S,R)	PTH ♦	(SST)	83970	(S,R)
AST	(SST)	84450	(S,R)	Electrolyte Panel	(SST)	80051	(S,R)	Renal Function Panel	(SST)	80069	(S,R)
Basic Metabolic Panel	(SST)	80048	(S,R)	FSH	(SST)	83001	(S,R)	Sodium	(SST)	84295	(S,R)
Beta HCG	(SST)	84702	(S,R)	Glucose	(SST)	82947	(S,R)	Testosterone	(SST)	84403	(S,R)
BUN	(SST)	84520	(S,R)	Hepatic Function Panel	(SST)	80076	(S,R)	Total Bilirubin	(SST)	82247	(S,R)
Calcium	(SST)	82310	(S,R)	HDL	(SST)	83718	(S,R)	Total Protein	(SST)	84155	(S,R)
CBC with Plt	(LT)	85027	(WB,R)	LH	(SST)	83002	(S,R)	Triglycerides	(SST)	84478	(S,R)
CBC with Plt & Diff	(LT)	85025	(WB,R)	Lipid Panel	(SST)	80061	(S,R)	TSH	(SST)	84443	(S,R)
Chloride	(SST)	82435	(S,R)	Magnesium	(SST)	83735	(S,R)	Unbound Testosterone	(SST)	84402	(S,R)
Cholesterol	(SST)	82465	(S,R)	Phosphorus	(SST)	84100	(S,R)	Uric Acid	(SST)	84550	(S,R)
CO <sub>2</sub>	(SST)	82374	(S,R)	Potassium	(SST)	84132	(S,R)				

**TUBE REQUIREMENTS:** SST-Serum Separator Tube LT-Lavender Top

**SPECIMEN REQUIREMENTS:** F-Frozen S-Serum R-Refrigerate U-Urine WB-Whole Blood  
♦ Must be processed within 48 hours of collection if not received frozen

**MicroCytePLUS®/Urine Cytology**

Test	Urine Collection Method	CPT
994 Hematuria Profile-Urine Cytology	Voided, Catheterized, Post-Cysto Void	88108, 88313, 81003, 82232, 82043, 84156
K600D Bladder Cancer FISH Pathodiagnostic	Voided, Catheterized, Post-Cysto Void, Wash (Bladder, Renal, Ureter)	88120
VU1D Bladder Cancer FISH/Cytology Pathodiagnostic Profile	Voided, Catheterized, Post-Cysto Void, Wash (Bladder, Renal, Ureter)	88112, 88120
VU4D Bladder Cancer FISH Reflex/Cytology Pathodiagnostic	Voided, Catheterized, Post-Cysto Void, Wash (Bladder, Renal, Ureter)	88112, if reflexed, 88120
VU3 Cytology Plus Monitoring Profile	Voided, Catheterized, Post-Cysto Void, Wash (Bladder, Renal, Ureter), Ileal Conduit/Neobladder	88112
VU6 Cytology Pap Stain Only	Voided, Catheterized, Post-Cysto Void, Wash (Bladder, Renal, Ureter), Ileal Conduit/Neobladder	88112

**EXPLANATION OF MicroCytePLUS®/URINE CYTOLOGY TESTING**

Hematuria Profile I - Urine Cytology for directing further evaluations of patients currently not monitored for TCC who present with hematuria or other signs of urinary tract or renal disease. Feulgen performed on all urine CYTOLOGY PROFILES.

Urine Volume – Provide a minimum of 50mL urine for optimum cellularity.  
Urine Viability – Hematuria to 5 days, Bladder Cancer FISH to 7 days, Cytology to 8 days.

Bladder Cancer FISH Cytology Pathodiagnostic Profile for therapeutic monitoring of patients with a history of TCC and for initial diagnosis of patients presenting with hematuria with suspicions of TCC.  
Bladder Cancer FISH Pathodiagnostic Test is Bladder Cancer FISH Assay, including diagnostic interpretation with clinical correlation by pathologist (MD).  
Bladder Cancer FISH results are intended for use as a method for monitoring for tumor recurrence in patients previously diagnosed with bladder cancer.  
Bladder Cancer FISH will not be performed on Ileal Conduit/Neobladder urine specimens.

**EXPLANATION OF REFLEX TESTING**

**Reflex Free PSA Testing**

Free PSA will be performed and billed if "Reflex Free PSA" is requested and the total PSA results fall within the requesting physician's previously defined parameters. The default range is 2-10 ng/ml.

**Quantitative Cystine**

When a qualitative cystine is positive, a quantitative cystine will be performed at an additional charge.

**Specimen Collection Information**

- \*Avoid submitting tissue specimens on fibrous materials such as gauze.
- \*After tissue has been obtained, place biopsy into 10% Formalin immediately. Do Not allow to air dry.
- \*All Urine Cytologies, FNA's and Fluid Aspirates must be submitted in the cytology alcohol fixative provided.
- \*Hematuria Profile Specimens must be collected in a preservative tablet kit.
- \*All 24 Hour Urine Specimens must be collected in a Dianon 24 Hour Urine Specimen Collection Kit (orange collection container) and submitted in the two vials provided with the kit.

**HELPFUL HINTS**

**24 Hour Urine Specimens** - Do Not Collect First Morning Void

**Ordering Kits**

Kit Orders may be placed through our Client Relations Department at 800-411-1839.

Please do not return unused histology vials or fixative bottles to our lab. Please dispose of unused histology vials in accordance with local laws and regulations regarding formalin disposal.

**DESCRIPTION OF PRIMARY LAB TESTING**

**AMA PANELS**

**Electrolyte Panel** 80051 - Sodium, Potassium, Chloride, Carbon Dioxide

**Lipid Panel** 80061@% - Chol@%, HDL@%, LDL (calculated)@%, Triglycerides@%

**Hepatic Function Panel** 80076 - Total Protein, Albumin, Total Bilirubin, Direct Bilirubin, Alkaline Phosphatase, SGOT (AST), SGPT (ALT)

**Basic Metabolic Panel** 80048 - Calcium, CO<sub>2</sub> (Carbon Dioxide), Chloride, Creatinine, Glucose@%, Potassium, Sodium, Urea Nitrogen (BUN)

**Renal Function Panel** 80069 - Albumin, Calcium, CO<sub>2</sub> (Carbon Dioxide), Chloride, Creatinine, Glucose@%, Phosphorus Serum, Potassium, Sodium, Urea Nitrogen (BUN)

**Comprehensive Metabolic Panel** 80053 - SGPT (ALT), Albumin, Total Bilirubin, Calcium, Chloride, Creatinine, Glucose@%, Alkaline Phosphatase, Potassium, Total Protein, Sodium, SGOT (AST), Urea Nitrogen (BUN), CO<sub>2</sub> (Carbon Dioxide)

**PROFILES**

**Hematology** 85027@ / 85025@ - CBC with PLT@, CBC with PLT and Diff@

**24 Hr Urine CPT Codes**

Ammonia 82140, Calcium 82340, Chloride 82436, Citrate 82507, Creatinine 82570, Magnesium 83735, Oxalate 83945, pH 83986, Phosphorus 84105, Potassium 84133, Qualitative Cystine 82127, Quantitative Cystine 82131, Sodium 84300, Sulfate 84392, Total Protein 84156, Uric Acid 84560

ProMark® is a registered trademark and service mark of Metamark Genetics, Inc. ConfirmMDx test performed and billed by MDxHealth® at Irvine, CA.

**Test Combination/Panel Policy**

Labcorp's policy is to provide physicians, in each instance, with flexibility to choose appropriate tests to assure that the convenience of ordering test combinations/panels do not distance physicians who wish to order a test combination/panel from making deliberate decisions regarding which tests are truly medically necessary. All the tests offered in test combinations/panels may be ordered individually using the Labcorp request form. Labcorp encourages clients to contact their local Labcorp representative or Labcorp location if the testing configurations shown here do not meet individual needs for any reason, or if some other combination of procedures is desired.

In an effort to keep our clients fully informed of the content, charges and coding of its test combinations/panels when billed to Medicare, we periodically send notices concerning customized test combinations/panels, as well as information regarding patient fees for all Labcorp services. We also welcome the opportunity to provide, on request, additional information in connection with our testing services and the manner in which they are billed to physicians, health care plans, and patients.

The CPT code(s) listed here are in accordance with the current edition of Current Procedural Terminology, a publication of the American Medical Association. CPT codes are provided here for the convenience of our clients; however, correct coding often varies from one carrier to another. Consequently, the codes presented here are intended as general guidelines and should not be used without confirming with the appropriate payor that their use is appropriate in each case. All laboratory procedures will be billed to third-party carriers (including Medicare and Medicaid) at fees billed to patients and in accordance with the specific CPT coding required by carrier. Microbiology CPT code(s) for additional procedures such as susceptibility testing, identification, serotyping, etc. will be billed in addition to the primary codes when appropriate. Labcorp will process the specimen for a Microbiology test based on source.