

CPU-to-CPU®

Technical Specification Guide

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Introduction

Exam*One's* CPU-to-CPU® system transmits laboratory results directly from Exam*One's* mainframe system to your system in a structured "raw data" format, rather than in a text report format. This raw data format has been Y2K compliant since version 6.03 of this technical document. The data is sent as a file containing multiple records for each applicant and company code, and when applicable, for multiple company codes. The flexible record format of this system allows testing information to be modified rapidly with very few changes for you and Exam*One*.

You can manipulate this data within your own automated or expert underwriting system to match applicant information to internal files, produce your own underwriting reports, or generate in-house statistics.

Appendix A contains sample reports of actual data for applicants generated by Exam*One* for client companies. These examples are provided in the Text File Format with its corresponding Raw Data File Format. Examples include:

- Standard UA
- Standard UA with UHIV
- Standard Oral Fluid Screen
- Standard Blood Profile & UA with Prostate Specific Antigen Report & Lipid Profile & Hepatitis Screen

Document Use

This document was written for our client's information systems personnel. It explains the system and how it may be used. Your application software should be capable of accepting all available results even if your company chooses not to perform one or more tests. Your system should be designed to accept any new Result Id without causing software failure. Exam*One* will notify clients in advance of any anticipated changes to the CPU-to-CPU® format. Every attempt will be made to give adequate lead-time for any changes. However, advance notification often is only a matter of weeks.

Exam*One* and EDI

Exam*One*, a Quest Diagnostics subsidiary is an active member of the Life/Annuity Task Group of the ASC X12N Insurance Subcommittee of the American National Standards Institute (ANSI). This group is chartered with developing Electronic Data Interchange (EDI) standardized messages for the sharing of information with our North American trading partners. The EDI format for ordering underwriting services was approved for use by the insurance underwriting industry in 1999 and is now available for use.

ANSI's long term strategy is to move EDI standardized messages to the International Standards that are set by the United Nations under UN/EDIFACT. This began in 1995. Exam*One* is dedicated to the efforts to standardize information exchange and will continue to be an active member of this task group. As EDI and UN/EDIFACT formatted messages are defined and approved for use, Exam*One* will support them for use by our clients.

Data Transmissions

Exam*One* transmits raw data files to hundreds of locations each day using a variety of transmission methods. We use a combination of synchronous and asynchronous modems at various speeds to ensure compatibility with your equipment. Our primary communication methods are listed in this section. Contact the Exam*One* Client Technical Support line at (800) 388-4675 for more information.

Transmissions via LabOne NETtm

Exam*One* has designed a communication network for transferring data between insurance companies and various information providers such as laboratories, paramedical, and inspection agencies. This network, referred to as LabOne NETtm, provides for automated transmission and receipt of our raw data. The primary LabOne NETtm distribution method is:

• **Dial-in to LabOne NET**tm: Exam*One* can provide you with password protected dial-in access or a direct Internet socket connection to our data transmission system where you may download your data.

Other Raw Data Transmission Methods

- FTP: Exam One can encrypt your data files using PGP encryption software and send the data to an FTP server over the Internet.
- FTPS/SFTP: Exam*One* supports transporting your data files using the latest in secure FTP protocols over the internet..

Transmission Windows

If necessary, Exam*One* can set up specific time windows in which your raw data transmissions will be performed. Contact Exam*One* for more details on scheduling.

Implementation Schedule

The following areas must be addressed before production implementation:

- Project Authorization;
- Communication hardware and software (including phone lines and modems);
- Client application software;
- Security requirements;
- Transmission testing between your site and Exam*One*;
- Final production authorization.

We will make every effort to conform to your implementation schedule. During testing and phased implementation, your current distribution methods will not be interrupted.

HIV and Sensitive Drugs Policy

Exam*One* is extremely concerned with the confidentiality of all applicant results. Our current policy is to not transmit sensitive (positive) drug or HIV results over any open communication lines. For complete confidentiality, applicant reports containing sensitive results will be sent by certified mail. However, with proper authorization from your company, sensitive results may be transmitted via the CPU-to-CPU® system. You will also have to modify your system in order to receive sensitive results. This functionality is located in Record 000, position 75. If you wish to receive sensitive results as raw data, please contact your sales representative for more information. See Appendix D for the form "Agreement to Provide Sensitive Test Results via Electronic Transmissions".

Your company has 3 options to choose from regarding the transmission of sensitive drug results.

- **Option 1:** This is the default option. If you elect not to receive sensitive results, the Special Status Indicator will be set to null. If an applicant has sensitive results, the whole applicant will be left off the transmission and the results will be mailed to you.
- **Option 2:** If you elect to receive sensitive results, the Special Status Indicator will be set to 'S' for those applicants containing sensitive results. You will receive both demographic information (Records 000-005) and results on all applicants.
- **Option 3:** If you choose not to have sensitive results electronically transmitted to you but want to receive notification within the data file, the comment "Results for this applicant are being mailed" will be displayed after the demographic records (000-005). The Special Status Indicator will be set to null. See the raw data example below.

File Specifications

A transmission file contains one or more company codes. Each company code contains one or more applicants. Each applicant has a minimum of seven 80-byte records generated. The maximum number of records depends on the number of tests requested for that applicant. The first three characters of each record indicate the record type. For a representation of a file, see **Figure 1** below.

000 APPLICANT #1 (FOR 1ST COMPANY CODE) DATA FOR 1ST COMPANY CODE POSSIBLY MORE APPLICANTS 998 COMPANY CODE 999 TRAILERS #1 000 APPLICANT #1 (FOR 2nd COMPANY CODE) DATA FOR 2ND COMPANY CODE POSSIBLY MORE APPLICANTS 998. COMPANY CODE TRAILERS #2 999

Figure 1: File Representation

Data for a single company code is made up of applicant records and company code records.

Applicant Records

- Each applicant must have record types 000-005 and a minimum of one record type from the 010-199 range.
- Record type 000 is used to separate all applicant records. Each applicant has one '000' record.
- Record types 001, 002, 003, 004 and 005 contain applicant demographics and/or specimen specific information. Each applicant receives one of each record type.

- Record types 010-099 contain specimen test results. These record types are numbered beginning with '010' and increase in increments of one. Each record contains a maximum of seven results. These record types contain Result Id range numbers 1000-8999.
- Record types 100-299 contain remarks. These record types are numbered beginning with '100' and increase in increments of one. You should design your system to accept several lines of sequence within each Remark Id. These record types contain Remark Id range numbers 9000-9899.

Company Code Records

- Record type 998 is reserved for future use with measures and ranges, and is not currently part of the transmission.
- Record type 999 contains record and applicant counts. One record exists per company code. This record type contains Total Id range numbers 9900-9999.

Additional File Information

Test results may be either numeric or alphanumeric. Numeric results contain only results with numbers. Alphanumeric results may contain spaces, numbers, letters and symbols. The CPU-to-CPU® system is designed for use with either field type.

The CPU-to-CPU® data file has been designed to allow for the addition of new test results without changing the file format. This allows you to receive the results of newly offered tests immediately upon request.

You are urged to design your systems to allow for this kind of flexibility. Your application software should be capable of accepting any new test results without causing a software failure (abort).

Suggestions for Design and Implementation

Underwriting Department Notification

A change in report formatting can cause problems for underwriters and medical personnel who use the test results. Notification of a change in the reporting structure before the change is implemented is imperative in achieving a smooth transition to a new report format.

Transmission Time

Data files will be available for transmission after 3:00 a.m. CST the following day.

Retransmissions

Two types of retransmissions may occur:

- A complete retransmission of **all** applicants for a particular processing cycle may be necessary, usually due to a loss of data on the receiving end of the transmission. Exam*One* retains several months of transmission data on-line to provide a timely and complete retransmission.
- A complete retransmission of **individual** applicant(s) may occur for various reasons but is usually due to a request from the client, or an update to the Exam*One* history files. When this type of retransmission occurs, all data for the requested/updated applicant(s) will be included in the next normally scheduled transmission.

For either type of retransmission, all available data for each applicant will be retransmitted. Because test results may be added, deleted or changed, your system should allow for total replacement of all applicant data or for the retention of complete multiple versions of all applicant data. Only the most current set of data should be considered complete, and reported or processed accordingly.

Report Identification

Appendix A contains sample applicant profiles generated by ExamOne. Whether you copy the ExamOne report format or design one of your own, to avoid confusion please indicate on the report that it was generated by your system.

Matching Criteria

An individual applicant can be uniquely identified in the Exam*One* database by using the combined data keys of "Date Performed" and "Control # (Lab Id)." However, this data will not be available to you until testing of the applicant is complete.

The following data keys may be used for matching applicant information:

• Ticket Number (Record Type 000)

The ticket number is preprinted on the Insurance Id Slip, which is sent to Exam*One* along with the specimens. The ticket number provides a unique key for each applicant **only if** the Id Slip has not been reproduced and/or submitted to Exam*One* more than once.

• Reference/Policy/Member Number (Record Type 000)

This data is obtained from the Id Slip under the Reference/Policy/Member Number field. Exam*One* provides this field for the client to record any of their own unique identifying codes. The accuracy and uniqueness of these codes is entirely dependent on the codes defined and used by your company and the diligence used when recording these codes on the Id Slip.

• Social Security Number (Record Type 000)

This number should not be used alone as a unique key because an applicant may be legitimately tested more than once, or it may not be included on the Id Slip.

• Applicant Name/Date of Birth/Sex (Record Type 001)

This combination of data fields may, in many cases, provide a unique key. However, it is not always reliable because of the commonality of names, birth dates, and sex.

The most reliable criteria for matching applicant information is a combination of several of the above data keys. The "best" combination of keys to use is determined, in part, by your company's requirements and the procedures you establish to ensure the accurate recording of key information.

Code Validation

The following Id ranges normally will not be changed: Data Id, Result Id, Remark Id, and Totals Id. However, specific Ids within these ranges can, and will be, added or deleted. Exam*One* will notify clients in advance of any anticipated changes. Advance notification often is only a matter of weeks. Therefore, your system design should be flexible, table-driven (if possible), and should avoid any validation that requires the existence of specific Ids. In addition, any new or unexpected Ids you receive should be handled routinely (in an exception manner) without causing a software failure (abort).

Record Descriptions

ABBREVIATION	DESCRIPTION
С	Constant – Contains a Exam <i>One</i>
	assigned value and should not be used
	for editing purposes.
N	Numeric – Contains only numbers.
AN	Alphanumeric – Contains spaces, letters,
	numbers, or symbols.
R	Required
0	Optional

Age Default: If AGE = 00, use 36.

Sex Default: If SEX is unknown (U), use MALE.

SSN Default: If SSN is blank, system defaults to spaces.

DOB Default: If DOB is blank, system defaults to spaces.

Demographic Information (000)

Position	Data Type	Length	Use	Field	Format/Values
01-03	N	3	R	Record Type	"000"
04-07	C	4	R	Certification Body	"HCFA"
08-17	С	10	R	Certification Number	"17D0648226"
18-25	N	8	R	Date Performed	CCYYMMDD
26-40	AN	15	R	Ticket Number	
41-65	AN	20	О	Reference/Policy/Member	
				Number	
66-74	N	9	О	SSN	
75	AN	1	О	Special Status Indicator	S = Sensitive; otherwise space
76-77		2		Spaces	
78-80	C	3	R	Identification Code	"HRL"

Demographic Information (001)

Position	Data Type	Length	Use	Field	Format/Values
01-03	N	3	R	Record Type	"001"
04-07	С	4	R	Data Id	"0100"
08-21	N	14	R	Control Number/Lab Id	ExamOne assigned
22-25	C	4	R	Data Id	"0125"
26-45	AN	20	R	Applicant Last Name	
46-49	С	4	R	Data Id	"0150"
50-60	AN	11	R	Applicant First Name	
61-64	С	4	R	Data Id	"0175"
65	AN	1	О	Applicant Middle Initial	
66-67	С	2	R	Data Id	"02"
68-75	N	8	R	Applicant Date of Birth	CCYYMMDD
76-79	С	4	R	Data Id	"0225"
80	AN	1	R	Sex of Applicant	"M" = Male
					"F" = Female
					"U" = Unknown

Demographic Information (002)

Position	Data Type	Length	Use	Field	Format/Values
01-03	N	3	R	Record Type	"002"
04-07	С	4	R	Data Id	"0250"
08-16		9		Spaces	
17-20	С	4	R	Data Id	"0275"
21-24	AN	4	R	Company Code	ExamOne assigned
25-28	С	4	R	Data Id	"0300"
29	AN	1	О	Life	"L"
30	AN	1	О	Health	"H"
31	AN	1	О	Disability	"D"
32	AN	1	О	Group	"G"
33	AN	1	О	Individual	"I"
34-37	С	4	R	Data Id	"0325"
38-47	N	10	О	Insurance Amount	Whole dollars
48-51	С	4	R	Data Id	"0350"
52-63	AN	12	О	Applicant City	
64-67	С	4	R	Data Id	"0375"
68-69	AN	2	О	Applicant State	
70-73	С	4	R	Data Id	"0400"
74-78	AN	5	О	Agency Code	
79-80	AN	2	0	Insurance Type Id	"CG"= Critical Illness – Group "CI" = Critical Illness – Individual "DG"= Disability – Group "DI" = Disability – Individual "HG"= Health – Group "HI"= Health – Individual "LG"= Life – Group "LI"= Life – Individual "MG"= Major Medical – Group "MI"= Major Medical – Individual "TG"= Long Term Care – Group "TI" = Long Term Care - Individual

Demographic Information (003)

Position	Data Type	Length	Use	Field	Format/Values
01-03	N	3	R	Record Type	"003"
04-07	С	4	R	Data Id	"0425"
08-21	AN	14	О	State/Agent	
22-25	С	4	R	Data Id	"0450"
26-39	AN	14	О	Examiner	
40-41	С	2	R	Data Id	"04"
42-49	N	8	О	*Date Collected (Urine)	CCYYMMDD
50-51	С	2	R	Data Id	"05"
52-59	N	8	О	Date of Last Meal	CCYYMMDD
60-63	С	4	R	Data Id	"0525"
64-68	AN	5	О	Time of Last Meal	HHMMX, where $x = \text{``A''}(AM)$ or "P" (PM)
69-72	С	4	R	Data Id	"0550"
73-80	N	8	О	*Date Collected (Serum)	CCYYMMDD

^{*}If Oral Fluid only specimen, date Oral Fluid specimen was collected will be in both Date Collected fields, otherwise, date Oral Fluid specimen was collected will be in the Date Collected (Serum) field.

Demographic Information (004)

Position	Data Type	Length	Use	Field	Format/Values
01-03	N	3	R	Record Type	"004"
04-07	C	4	R	Data Id	"0575"
08-12	AN	5	О	*Time Collected	HHMMX, where $x = \text{``A''}(AM) \text{ or ``P''}(PM)$
13-16	С	4	R	Data Id	"0600"
17	AN	1	R	Signature Present	"Y" = Yes
					"N" = No
18-19	С	2	R	Data Id	"06"
20-27	N	8	R	Date Completed	CCYYMMDD
28-31	C	4	R	Data Id	"0650"
32-46	AN	15	О	Agent City	
47-50	C	4	R	Data Id	"0675"
51-52	AN	2	О	Agent State	
53-56	С	4	R	Data Id	"0680"
57-65	AN	9	О	Agent Zip Code	
66-69	C	4	R	Data Id	"0690"
70-73	AN	4	О	Examiner Company Id	
74-77	C	4	R	Data Id	"0695"
78-79	AN	2	О	Examiner State	
80		1		Space	

^{*}Time is supplied by the examiner and is captured from the Id slip. No validation is performed.

Note: Date Completed (Record 004, position 20) reflects the date assigned or updated by Exam*One's* automated laboratory system each time an applicant's record is modified. The Date Completed field will match the Date Performed (Record 000, position 18) the first time an applicant's results are transmitted. Any subsequent transmission of an applicant's record includes the entire record with requested changes, and reflects the date of the change in the Date Completed field, but preserves the original Date Performed. By comparing the two dates, modified records may be easily identified and processed. To ensure data consistency, all previous information on the applicant should be replaced with the updated record.

Demographic Information (005)

Position	Data Type	Length	Use	Field	Format/Values
01-03	N	3	R	Record Type	"005"
04-07	С	4	R	Data Id	"0700"
08-42	AN	35	О	Applicant Address	
43-46	С	4	R	Data Id	"0725"
47-55	AN	9	О	Applicant Zip Code	
56-59	С	4	R	Data Id	"0750"
60-79	AN	20	О	Agent/Agency Name	
80		1		Space	

Test Results (010-099)

⇒ Every applicant has at least one result or remark record. Result record types range from 010-099. Each result record contains a maximum of seven results. Each result is identified by a unique Result Id. (See *Result Ids* in Appendix B)

Position	Data Type	Length	Use	Field	Format/Values
01-03	N	3	О	Record Type	"010-099"
04-07	N	4	О	Result Id#	"1000-8999"
08-14	AN	7	О	*Result Value	
15-18	N	4	О	Result Id #	"1000-8999"
19-25	AN	7	О	*Result Value	
26-29	N	4	О	Result Id#	"1000-8999"
30-36	AN	7	О	*Result Value	
37-40	N	4	О	Result Id #	"1000-8999"
41-47	AN	7	О	*Result Value	
48-51	N	4	О	Result Id #	"1000-8999"
52-58	AN	7	О	*Result Value	
59-62	N	4	О	Result Id#	"1000-8999"
63-69	AN	7	О	*Result Value	
70-73	N	4	О	Result Id #	"1000-8999"
74-80	AN	7	О	*Result Value	

^{*}If the result value is numeric, the format is 7 positions with 3 positions to the right of the (implied) decimal point (9999.999). If the result value is alphanumeric, the format is left justified, space-filled (XXXXXXX) and represents a valid result or replacement remark.

Test Remarks (100-299)

- ⇒ Each record contains one test remark identified by a remark identification number. The remark can be associated with one or more Result Ids or a group of Result Ids. (See *Remark Ids and Associated Result Ids* in Appendix C)
- ⇒ Occasionally it becomes necessary to provide the client with more information about a test than can be accommodated in one Remarks Record. In this instance, an additional Remarks Record(s) will be created that contain the same Remark Id #. See example below.

Position	Data Type	Length	Use	Field	Format/Values
01-03	N	3	R	Record Type	"100-299"
04-07	N	4	R	Remark Id #	"9000-9899"
08-80	AN	70	О	Comment	

Raw Data Example:

```
1009700 COMPANY NOTIFIED OF CONTAINER IDENTIFICATION PRIOR TO UPDATE. ===> Remark 1
1019850 ADULTERANT TESTS WITHIN NORMAL LIMITS ==> Remark 2
1029800 THE URINE HIV-1 TEST MAY MISS 1 TO 2% OF ALL TRUE POSITIVES. THE ==> Remark 3
1039800 TEST MAY GENERATE UP TO 6% FALSE POSITIVES WHEN TESTING SUBJECTS AT
1049800 HIGH RISK FOR HIV-1 INFECTION AND 10% FALSE POSITIVES FOR INDIVIDUALS
1059800 WITH OTHER MEDICAL CONDITIONS.
```

Ranges and Measures (998)

⇒ Ranges may be included in a future version of the CPU-to-CPU® system. When this is added, the 998 records will include Result Ids with the measures and ranges in effect at the same time of transmission.

Position	Data Type	Length	Use	Field	Format/Values

Record Counts (999)

⇒ This record is included as the last entry for each company's data within the file. The record indicates the total number of applicants and the total number of records for an individual company code.

Position	Data Type	Length	Use	Field	Format/Values
01-03	N	3	R	Record Type	"999"
04-07	С	4	R	Total Id	"9925"
08-17	N	10	R	Number of Applicants	Per company code
18-21	С	4	R	Total Id	"9950"
22-31	N	10	R	Number of Records	Per company code
32-80		49		Spaces	

Appendix A – Applicant Report Examples

- > Standard UA
- > Standard UA with UHIV
- > Standard Oral Fluid Screen
- ➤ Standard Blood Profile & UA with Prostate Specific Antigen Report & Lipid Profile & Hepatitis Screen

Standard UA - Text File Format

LABONE, INC. CLIA NO.: 17D0648226 CAP ACCREDITATION NO.: 28646-01

10101 RENNER BLVD. LENEXA KS, 66219

MED. INS. DIRECTOR: RICHARD BRAUN, MD LAB DIRECTOR: PATRICK JAMES, MD

NAME: MARY B. AMMONS

DOB/SEX/ST: 09/02/1960 F MO

AGENT/AGENCY: XX/

EXAMINER: XX/ABC EXAM TICKET NUMBER: 0040522072

INS TYPE/AMT: IND LIFE/\$ 1,000,000

QUALITY LIFE INSURANCE DATE PERFORMED: 09/24/2001 DCL 38925062

10101 RENNER BLVD INSURANCE KEY:

LENEXA, KS 66219 D/T LAST MEAL: NOT GIVEN ATTN:LEACY BUSSELL D/T COLLECTED: 09/19/2001

MEDICAL DIRECTOR

SOC SEC NO:

-SPECIAL TESTS/DRUGS	5-	-URINALYSIS-			-REFERENC	CE RANGE-
DIURETIC AGTS (DIU)	NEG	GLUCOSE	(GM%)	NEG		NEGATIVE
		PROTEIN	(MG%)	<10	0	- 30
BETA BLOCKERS (BAB)	NEG	LEUKOCYTE SCREEN		NEG		NEGATIVE
		HEMOGLOBIN SCREEN		NEG		NEGATIVE
COT(NIC) (MCG/ML)	NEG	WHITE BLOOD CELLS (/	/HPF)	NP	0	- 9
		RED BLOOD CELLS (/	/HPF)	NP	0	- 4
COCAINE	POS	GRANULAR CASTS (/40)LPF)	NP		0
		HYALINE CASTS (/40)LPF)	NP	0	- 10
		SPECIFIC GRAVITY		NP	1.003	- 1.035
		URINE TEMP(FAHRENHE	EIT)	98.0	90.5	- 99.6

ADULTERANT TESTS WITHIN NORMAL LIMITS

Corresponding Raw Data File Format

 0000HCFA17D0648226200109240040522072
 S HRL

 0010100000000389250620125AMMONS
 0150MARY
 0175B02196009020225F

 0020250
 0275DCL
 0300L
 1032500010000000350KC
 0375M00400

 0020250
 0275DCL
 0300L
 I03250001000000350KC
 0375M00400

 0030425XX
 0450ABC
 EXAM
 042001091905
 0525
 055020010919

 0040575
 0600Y06200109240650
 0675XX0680
 0690GLF
 0695XX

0050700 0725 0750

010525000090005050NEG 4975NEG 5000NEG 5225NEG 6005009800050250001000

01160150038100602000051007125NEG 7130NEG 1009850ADULTERANT TESTS WITHIN NORMAL LIMITS

Standard UA with UHIV - Text File Format

LABONE, INC. CLIA NO.: 17D0648226 CAP ACCREDITATION NO.: 28646-01

10101 RENNER BLVD. LENEXA KS, 66219

MED. INS. DIRECTOR: RICHARD BRAUN, MD LAB DIRECTOR: PATRICK JAMES, MD

NAME: JOAN THOMPSON

DOB/SEX/ST: 06/20/1978 F XX

AGENT/AGENCY: XX/

EXAMINER: XX/ABC EXAM TICKET NUMBER: 0040484715

INS TYPE/AMT: IND LIFE/NOT GIVEN

QUALITY LIFE INSURANCE DATE PERFORMED:08/03/2001 DCL 38249881

10101 RENNER BLVD INSURANCE KEY:

LENEXA, KS 66219 D/T LAST MEAL: NOT GIVEN ATTN:LEACY BUSSELL D/T COLLECTED: 08/01/2001

MEDICAL DIRECTOR

SOC SEC NO:

_	-URINALYSIS-			-REFERENCI	E RANGE-
NEG	GLUCOSE	(GM%)	NEG	1	NEGATIVE
	PROTEIN	(MG%)	17	0 -	- 30
NEG	LEUKOCYTE SCREEN		NEG]	NEGATIVE
NEG	HEMOGLOBIN SCREEN		NEG]	NEGATIVE
NEG	WHITE BLOOD CELLS	(/HPF)	NP	0 -	- 9
	RED BLOOD CELLS	(/HPF)	NP	0 -	- 4
NEG	GRANULAR CASTS (/4	OLPF)	NP	(0
	HYALINE CASTS (/4	OLPF)	NP	0 -	- 10
	SPECIFIC GRAVITY		NP	1.003	- 1.035
	URINE TEMP (FAHREN	HEIT)	98.0	90.5	- 99.6
	NEG NEG NEG NEG	NEG GLUCOSE PROTEIN NEG LEUKOCYTE SCREEN NEG HEMOGLOBIN SCREEN NEG WHITE BLOOD CELLS RED BLOOD CELLS NEG GRANULAR CASTS (/4 HYALINE CASTS (/4 SPECIFIC GRAVITY	NEG GLUCOSE (GM%) PROTEIN (MG%) NEG LEUKOCYTE SCREEN NEG HEMOGLOBIN SCREEN NEG WHITE BLOOD CELLS (/HPF) RED BLOOD CELLS (/HPF) NEG GRANULAR CASTS (/40LPF) HYALINE CASTS (/40LPF)	NEG GLUCOSE (GM%) NEG PROTEIN (MG%) 17 NEG LEUKOCYTE SCREEN NEG NEG HEMOGLOBIN SCREEN NEG WHITE BLOOD CELLS(/HPF) NP RED BLOOD CELLS (/HPF) NP NEG GRANULAR CASTS (/40LPF) NP HYALINE CASTS (/40LPF) NP SPECIFIC GRAVITY NP	NEG GLUCOSE (GM%) NEG PROTEIN (MG%) 17 0 NEG LEUKOCYTE SCREEN NEG NEG HEMOGLOBIN SCREEN NEG NEG WHITE BLOOD CELLS(/HPF) NP 0 RED BLOOD CELLS (/HPF) NP 0 NEG GRANULAR CASTS (/40LPF) NP HYALINE CASTS (/40LPF) NP SPECIFIC GRAVITY NP 1.003

URINE HIV-1 ANTIBODY SCREEN: NON-REACTIVE

THE URINE HIV-1 TEST MAY MISS 1 TO 2% OF ALL TRUE POSITIVES. THE TEST MAY GENERATE UP TO 6% FALSE POSITIVES WHEN TESTING SUBJECTS AT HIGH RISK FOR HIV-1 INFECTION AND 10% FALSE POSITIVES FOR INDIVIDUALS WITH OTHER MEDICAL CONDITIONS.

		URINE ADULTER	ANT RESULTS		
DETERMINAT	CION	ABNORMAL	NORMAL	NORMAL I	IMITS
CREATININE	(MG/DL)		154.2	>	5.0
PH			7.7	4.0 -	8.7
URINE TEMPERAT	URE		98.0	90.5 -	99.6

Corresponding Raw Data File Format

000HCFA17D0648226200108030040484715			HRL
001010000000382498810125THOMPSON	0150JOAN	0175 0219	7806200225F
0020250 0275DCL 0300L I0325000	0000000350	0375XX	0400
0030425XX	42001080105	0525 0	55020010801
0040575 0600Y06200108030650	0675XX068	0690	GLF 0695XX
0050700	0725	0750	
010525000170005050NEG 4975NEG 5000	NEG 5225NEG	60050098000	50250000000
0116000NEG 60150154200602000077005235	NEG 7125NEG	7130NEG	6007NEG
1009850ADULTERANT TESTS WITHIN NORMAL LI	MITS		
1029800THE URINE HIV-1 TEST MAY MISS 1 T	O 2% OF ALL TRU	E POSITIVES.	THE
1039800TEST MAY GENERATE UP TO 6% FALSE	POSITIVES WHEN '	resting subject	TS AT
1049800HIGH RISK FOR HIV-1 INFECTION AND	10% FALSE POSI	TIVES FOR INDI	VIDUALS
1059800WITH OTHER MEDICAL CONDITIONS.			

Standard Oral Fluid – Text File Format

LABONE, INC. CLIA NO.: 17D0648226 CAP ACCREDITATION NO.: 28646-01

10101 RENNER BLVD. LENEXA KS, 66219

MED. INS. DIRECTOR: RICHARD BRAUN, MD LAB DIRECTOR: PATRICK JAMES, MD

ORAL FLUID SCREEN

NAME: FLORA NELLS

DOB/SEX/ST: 01/10/1968 F XX

AGENT/AGENCY: XX/

EXAMINER: XX/ABC EXAM TICKET NUMBER: 0040487427

INS TYPE/AMT: IND LIFE/NOT GIVEN

QUALITY LIFE INSURANCE DATE PERFORMED:09/10/2001 DCL 38724197

INSURANCE KEY: 10101 RENNER BLVD

D/T LAST MEAL: NOT GIVEN LENEXA, KS 66219 ATTN:LEACY BUSSELL D/T COLLECTED: 09/05/2001

MEDICAL DIRECTOR

SOC SEC NO:

ORAL FLUID HIV ANTIBODY STATUS: NEGATIVE

----- ORAL FLUID DRUGS ------

DETERMINATION RESULT COCAINE COTININE (NIC) NEG

Corresponding Raw Data File Format

000HCFA17D0648226200109100040487427

0050700 0725 0750

0106525Y 6550POS 6555NEG 6520NEG

9999925000000000199500000000008

Standard Blood Profile & UA with Prostate Specific Antigen Report & Lipid Profile & Hepatitis Screen - Text File Format

CLIA NO.: 17D0648226 CAP ACCREDITATION NO.: 28646-01 LABONE, INC.

10101 RENNER BLVD. LENEXA KS, 66219

MED. INS. DIRECTOR: RICHARD BRAUN, MD LAB DIRECTOR: PATRICK JAMES, MD

NAME: STEPHEN J. TIMBLE

DOB/SEX/ST: 02/09/1960 M MO

AGENT/AGENCY: XX/ EXAMINER: XX/

TICKET NUMBER: 0040521955

INS TYPE/AMT: IND LIFE/\$ 1,000,000

QUALITY LIFE INSURANCE DATE PERFORMED:07/02/2001 DCL 37798888

10101 RENNER BLVD INSURANCE KEY:

LENEXA, KS 66219 D/T LAST MEAL: NOT GIVEN

ATTN:LEACY BUSSELL D/T COLLECTED: 06/28/2001 D/T COLLECTED: 06/28/2 SERUM APPEAR: NORMAL MEDICAL DIRECTOR

SOC SEC NO: ZIP:64113

		500 5E0					·
		FULL BLOOD CHEM			USUAL RA	CL	INICAL
		-ABNORMAL-					
	(MG/DL)		1	08	60		
FRUCTOSAMINE			1	.5	1.2	-	2.0
		NOT PERFORMED			3.0	-	6.0
	(MG/DL)			24	9		27
CREATININE		2.4 H	1		0.7		
ALK. PHOS.					30		
BILI. TOT.	(MG/DL)	66 Н	0	. 7	0.2		
					-		33
		64 H					45
		30			-) –	65
TOT. PROTEIN	(G/DL)	4.4 L			6.1	-	8.2
ALBUMIN					3.8		
GLOBULIN	(G/DL)	1.7 L			2.1	-	3.9
CHOLESTEROL		126 L			140		
HDL CHOLESTEROL	(MG/DL)			58			80
LDL (CALCULATED)	(MG/DL)				0	-	129
CHOL/HDL CHOL RAT	TIO		2	.2		<	
LDL/HDL RATIO		0.74 L			0.9	-	5.3
TRIGLYCERIDES				26		-	150
		PRESENT TO PERFORI					
-SPECIAL TESTS/DE	RUGS-	-URINALYSIS-			-REFERENC	CE	RANGE-
DIURETIC AGTS (DIU	J) NEG	GLUCOSE	(GM%)	0.25		NE	GATIVE
		PROTEIN	(MG%)	15	0	-	30
BETA BLOCKERS (BAR	B) NEG	PROTEIN LEUKOCYTE SCREEN HEMOGLOBIN SCREEN		NEG		NE	GATIVE
		HEMOGLOBIN SCREEN		NEG		NE	GATIVE
COT(NIC) (MCG/MI	L) NEG	WHITE BLOOD CELLS	(/HPF)	NP	0	_	9
		RED BLOOD CELLS		NP	0	_	4
COCAINE	NEG	GRANULAR CASTS (/		NP		0	
		HYALINE CASTS (/			0	_	10
					1.003	_	1.035
		URINE TEMP(FAHREN	HEIT) N	OT GIVEN	90.5	_	99.6
		ADULTERANT TE					

LABONE, INC. CLIA NO.: 17D0648226 CAP ACCREDITATION NO.: 28646-01

10101 RENNER BLVD. LENEXA KS, 66219

MED. INS. DIRECTOR: RICHARD BRAUN, MD LAB DIRECTOR: PATRICK JAMES, MD

PROSTATE SPECIFIC ANTIGEN REPORT

NAME: STEPHEN J. TIMBLE

DOB/SEX/ST: 02/09/1960 M MO

AGENT/AGENCY: XX/ EXAMINER: XX/

TICKET NUMBER: 0040521955

INS TYPE/AMT: IND LIFE/\$ 1,000,000

QUALITY LIFE INSURANCE DATE PERFORMED:07/02/2001 DCL 37798888

10101 RENNER BLVD INSURANCE KEY:

LENEXA, KS 66219 D/T LAST MEAL: NOT GIVEN
ATTN:LEACY BUSSELL D/T COLLECTED: 06/28/2001
MEDICAL DIRECTOR SERUM APPEAR: NORMAL

SOC SEC NO: ZIP:64113

PROSTATE SPECIFIC ANTIGEN RESULTS

DETERMINAT	ION	OUT OF RANGE	NORMAL	USUAL RANGE
PSA	(NG/ML)	9.84		< 4.01
DEBCENT ERFE DON	121	OUDNITTY NOT SUI	FFICIENT	

<10 %FREE PSA INCREASED RISK OF CANCER
10-25 %FREE PSA INTERMEDIATE RISK OF CANCER
>25 %FREE PSA DECREASED RISK OF CANCER

THE PERCENT FREE PSA IS INVERSELY PROPORTIONAL TO

THE RISK OF PROSTATE CANCER.

PERCENT FREE PSA IS NOT ABSOLUTE EVIDENCE OF MALIGNANCY. PERCENT FREE PSA IS PERFORMED USING THE BECKMAN COULTER ACCESS II. VALUES OBTAINED WITH DIFFERENT ASSAY METHODS MAY NOT BE INTERCHANGEABLE.

LIPID PROFILE

PROPOSED INSURED: STEPHEN J. TIMBLE

DATE OF BIRTH: 02/09/1960

SEX: M

DATE PERFORMED: 07/02/2001

BELOW ARE SOME OF THE RESULTS OF THE BLOOD TEST RECENTLY PERFORMED IN CONJUNCTION WITH YOUR APPLICATION FOR INSURANCE. IF YOU HAVE ANY QUESTIONS REGARDING THESE RESULTS, PLEASE CONTACT YOUR PERSONAL PHYSICIAN.

DETERMINATION	RESULT	LT REFERENCE RANGE*				
TOTAL CHOLESTEROL	126 (MG/DL)		DESIRABLE BORDERLINE RISK HIGH RISK			
HDL	58 (MG/DL)	36 OR GREATER				
TOTAL CHOL./HDL RATIO	2.2	4.5 OR LESS				
LDL (CALCULATED) **	42 (MG/DL)	130 - 159	DESIRABLE BORDERLINE RISK HIGH RISK			
LDL/HDL RATIO**	0.74	LESS THAN 4 LESS THAN 3	MEN WOMEN			
TRIGLYCERIDES**	126 (MG/DL)	250 OR LESS				

^{*} THESE ARE THE STANDARDIZED RANGES OF THE NATIONAL CHOLESTEROL EDUCATION PROGRAM, THE AMERICAN HEART ASSOCIATION AND THE UNIVERSITY OF KANSAS MEDICAL CENTER. RANGES USED BY YOUR INSURER FOR UNDERWRITING MAY VARY.

^{**} THESE RESULTS MAY BE AFFECTED BY YOUR FASTING STATE AT THE TIME OF THE BLOOD DRAW. IF THE TRIGLYCERIDE VALUE IS GREATER THAN 400 MG/DL, THE LDL CALCULATION WILL NOT BE REPORTED.

LABONE, INC. CLIA NO.: 17D0648226 CAP ACCREDITATION NO.: 28646-01

10101 RENNER BLVD. LENEXA KS, 66219

MED. INS. DIRECTOR: RICHARD BRAUN, MD LAB DIRECTOR: PATRICK JAMES, MD

NAME: STEPHEN J. TIMBLE

DOB/SEX/ST: 02/09/1960 M MO

AGENT/AGENCY: XX/ EXAMINER: XX/

TICKET NUMBER: 0040521955

INS TYPE/AMT: IND LIFE/\$ 1,000,000

QUALITY LIFE INSURANCE DATE PERFORMED:07/02/2001 DCL 37798888

10101 RENNER BLVD INSURANCE KEY:

LENEXA, KS 66219 D/T LAST MEAL: NOT GIVEN
ATTN:LEACY BUSSELL D/T COLLECTED: 06/28/2001
MEDICAL DIRECTOR SERUM APPEAR: NORMAL

SOC SEC NO: ZIP:64113

HEPATITIS TEST -ABNORMAL- -NORMAL- --- RANGE ----

HBSAG NEGATIVE NEGATIVE HEPATITIS C Ab NEGATIVE NEGATIVE NEGATIVE

Corresponding Raw Data File Format

0020250 0275DCL 0300L 103250001000000350KANSAS CITY 0375M00400

0050700 072564113 0750

0101250000270012750100000130000070013250024000135001260001375000240014000030000
01114500066000147500640001500000440015750001700152501260001560000150014250108000
012160000580001625000220016300042000163500007402100NP 2685000984052500015000
013505000002504975NEG 5000NEG 5225NEG 6005NG 5025000000060150154900
01460200006700138001610007125NEG 7130NEG 4315NEG 4365NEG 2687QNS

1009100SPECIMEN WAS NOT PRESENT TO PERFORM REQUESTED GLYCO

1019850ADULTERANT TESTS WITHIN NORMAL LIMITS

1029300NORMAL

Appendix B

➤ Result Ids

Result Ids

SPEC TYPE = SPECIMEN	DESCRIPTION
В	Blood
U	Urine
S	Oral Fluid
D	Dried Blood Spot

Result Id	Remark Id	Test Description	Spec Type	Gender	Age	Reference Range	Unit of Measure	Decimal Position	Reference
1050	9000	CBC – HCT	В	F M/Unk	All	37-47 42-52	%	999.9 or ZZ9.9	
1075	9000	CBC – HGB	В	F M/Unk	All	12-16 14-18	g/dL	999.9 or ZZ9.9	
1100	9000	CBC – MCH	В	All	All	27.0-31.0	pg	999.9 or ZZ9.9	
1125	9000	CBC - MCHC	В	All	All	33.0-37.0	%	999.9 or ZZ9.9	
1150	9000	CBC – MCV	В	All	All	81.0-99.0	μ^3	999.9 or ZZ9.9	
1175	9000	CBC – PLT	В	All	All	130-400	x1,000	9999 or ZZZ9	
1200	9000	CBC – RBC	В	F M/Unk	All	4.2-5.4 4.6-6.2	x1,000,000	99.99 or Z9.99	
1225	9000	CBC – WBC	В	All	All	4.8-10.8	x1,000	999.9 or ZZ9.9	
1250	9100	Chemistry – Albumin	В	All	All	3.8-5.2	g/dL	9.9 or Z.9	
1251	9100	Chemistry – Albumin/Globulin Ratio	В	All	All	1.0 – 2.5	g/dL	99.9 or Z9.9	
1275	9100	Chemistry – Alkaline Phosphatase	В	All	0-18 19+	30-200 30-125	U/L	9999 or ZZZ9	
1300	9100	Chemistry – Total Bilirubin	В	F M/Unk	All	0.2-1.2 0.2-1.5	mg/dL	99.9 or Z9.9	
1325	9100	Chemistry – BUN	В	F F M/Unk M/Unk	0-50 51+ 0-50 51+	7-22 9-26 9-25 9-27	mg/dL	999, ZZ9 or ZZZ	
1330	9100	Chemistry-Pro-BNP	В	All	0 - 74 75+	0 – 124 0 – 449	pg/mL	9999, ZZZ9	
1350	9100	Chemistry – Total Cholesterol	В	All	All	140-199	mg/dL	9999 or ZZZ9	
1375	9100	Chemistry – Creatinine	В	F M/Unk	All	0.6-1.3 0.7-1.5	mg/dL	99.9 or Z9.9	
1380	9100	Chemistry – LDH	В	All	All	100-280	U/L	9999 or ZZZ9	
1400	9100	Chemistry – GGT	В	F M/Unk	All	0-45 0-65	U/L	9999 or ZZZ9	
1425	9100	Chemistry - Glucose	В	All	All	60-109	mg/dL	999, ZZ9 or ZZZ	
1450	9100	Chemistry – AST	В	All	All	0-33	U/L	9999 or ZZZ9	
1475	9100	Chemistry – ALT	В	All	All	0-45	U/L	9999 or ZZZ9	
1500	9100	Chemistry – Total Protein	В	All	All	6.1-8.2	g/dL	99.9 or Z9.9	

							Updated June 2023			
Result Id	Remark Id	Test Description	Spec Type	Gender	Age	Reference Range	Unit of Measure	Decimal Position	Reference	
1525	9100	Chemistry – Triglycerides	В	All	All	0-150	mg/dL	9999 or ZZZ9		
1560	9100	Chemistry – Fructosamine	В	All	All	1.2-2.0	mmol/L	99.9 or Z9.9		
1575	9100	Chemistry – Globulin (Total Protein minus albumin)	В	All	All	1.9-3.7	g/dL	99.9 or Z9.9		
1600	9100	Chemistry – HDL Cholesterol	В	F M/Unk	All	35-100 35-80	mg/dL	9999 or ZZZ9		
1625	9100	Chemistry – Cholesterol/ HDL Cholesterol Ratio	В	All	All	< 5.0		999.9 or ZZ9.9		
1630	9100	Chemistry – LDL Cholesterol	В	All	All	0-129	mg/dL	9999 or ZZZ9		
1635	9100	Chemistry – LDL/HDL Cholesterol Ratio	В	F M/Unk	All	0.6-4.3 0.9-5.3		999.99 or ZZ9.99		
1650	9650	Differential – Atypical Lymph	В	All	All			Alpha X(7)		
1675	9650	Differential – Band	В	All	All	0-7	%	999, ZZ9, or ZZZ		
1700	9650	Differential – Basophil	В	All	All	0-1	%	999, ZZ9 or ZZZ		
1725	9650	Differential – Blast	В	All	All	0-0	%	999, ZZ9 or ZZZ		
1750	9650	Differential – Eosinophil	В	All	All	1-5	%	999, ZZ9 or ZZZ		
1775	9650	Differential – Lymphocyte	В	All	All	24-46	%	999, ZZ9 or ZZZ		
1800	9650	Differential – Metacyte	В	All	All	0-0	%	999, ZZ9 or ZZZ		
1825	9650	Differential – Monocyte	В	All	All	1-10	%	999, ZZ9 or ZZZ		
1850	9650	Differential – Myelocyte	В	All	All	0-0	%	999, ZZ9 or ZZZ		
1875	9650	Differential –	В	All	All	0-0	%	999, ZZ9 or ZZZ		
1900	9650	Promyelocyte Differential – Segmented Neutrophil	В	All	All	35-66	%	999, ZZ9 or ZZZ		
2100	9100	Hemoglobin A1C	В	All	All	3.0-6.0	%	99.9 or Z9.9		
2125	9200	Serum HIV Interpretation	В	All	All	Neg		Alpha X(7)		
2680	9200	Beta – 2 Microglobulin	В	All	All	00.00 – 2.51	Mg/L	99.99 or Z9.9Z		
2685	9750	PSA – Prostate Specific Antigen	В	All	All	0.00-4.00	ng/mL	999.99 or ZZZ9.99		
2705	9100	Ethyl Alcohol	В	All	All	0 – 9	mg/dL	999, ZZ9		
2720	9890	CDT - Carbohydrate Deficient Transferrin	В	All	All	Neg		Alpha X(7)		
2722	9890	CDT – Quantitative	В	All	All	Neg		9.9 or Z9.9		
3095	9100	Cystatin C	В	F F F F F	0-18 19-29 30-39 40-49 50-59 60-69 70-79	0.52 - 1.19 $0.52 - 1.28$ $0.52 - 1.24$ $0.52 - 1.21$ $0.52 - 1.17$ $0.52 - 1.14$ $0.52 - 1.10$	mg/L	99.99 or Z9.99		
				F	80-89	0.52 - 1.10 $0.52 - 1.07$				

	Updated June 2023								2023
Result	Remark	Test Description	Spec	Gender	Age	Reference Range	Unit of	Decimal	Reference
Id	Id		Type				Measure	Position	
				F	90+	Not Available			
				M/Unk	0-18	0.52 - 1.19			
				M/Unk	19-29	0.52 - 1.35			
				M/Unk	30-39	0.52 - 1.31			
				M/Unk	40-49	0.52 - 1.27			
				M/Unk	50-59	0.52 - 1.27 $0.52 - 1.23$			
					60-69	0.52 - 1.20			
				M/Unk		0.52 - 1.20 $0.52 - 1.16$			
				M/Unk	70-79				
				M/Unk	80-89	0.52 - 1.13			
				M/Unk	90+	Not Available			
3135	9375	Dried Blood HIV Interpretation	D	All	All	Neg		Alpha X(7)	
22004	0.400	Toxicology-6-		4.11	4 11	0 = Negative	/ T	0 7	
3289^	9400	Monoacetylmorphine	U	All	All	1 = Positive	ng/mL	9 or Z	
3340	9175	Dried Blood Nicotine	D	All	All	Neg		Alpha X(7)	
3340	91/3		D	All	AII	neg			
3350	9175	Dried Blood – Total	D	All	All	0-199*	mg/dL	9999 or	
		Cholesterol				0 277	8	ZZZ9	
3664	9100	Vary Lovy Dansity Linid	В	All	All	1 - 40	ma/Dl	9999 or	
3004	J100	Very Low Density Lipid	Б	AII	AII	1 – 40	mg/Dl	ZZZ9	
					1	0.7-1.9			
			_		2-7	0.8-2.3		99.9 or	
3693	9130	FT4 – Free Thyroxine	В	All	8-20	0.6-2.0	ng/dL	Z9.9	
					21+	0.8-1.8		27.9	
					1-9*	127-221			
3701	9130	T-3 Total	В	All	10-13	123-211	ng/dL	999.9 or	
3701	7150	1 5 10111		7 111	14-18	97-186	ng/uL	ZZ9.9	
					19+	97-219			
		TOLL TI '1			1	0.8-8.2		000.00	
3703	9130	TSH – Thyroid	В	All	2-20	0.7-5.7	μIU/mL	999.99 or	
3703	7150	Stimulating Hormone		7 111	21+	0.35 - 5.50	μιοππΕ	ZZ9.99	
								9999.9,	
3950	9750	Alpha –Fetoprotein	В	All	All	0 - 6.1	ng/mL		
						G 1		ZZZ9.9	
		CEA – Carcinoembryonic	_			Smoker:			
3952	9750	Antigen	В	All	All	0.0-5.0	ng/mL	ZZZ9.9	
		Anugen				Non-Smoker: 0.0-2.5			
2050	07.50	D. F. D.C.	Б	. 11	. 11	. 0.5	0./	ZZ9	
3959	9750	Percent Free PSA	В	All	All	>25	%		
		Toxicology –				0 = Negative		1	
4010^	9400		U	All	All	1 = Positive		9	
		Benzodiazepine						+	
4012^	9400	Toxicology – Barbiturate	U	All	All	0 = Negative		9	
	2.00	<u>.</u>				1 = Positive			
4014^	9400	Toxicology – Methadone	T T	A 11	A 11	0 = Negative		9	
4014	9400	Qualitative CLS	U	All	All	1 = Positive		9	
			_					99.9 or	
4144	9130	T-3 Uptake	В	All	All	22.0-35.0*	%	Z9.9	
4145	9130	Free T3	В	All	All	2.3 – 4.2	PG/mL	999.9	
7143	713U	1100 13	D	All			1 U/IIIL	777.7	
					1-4	7.2-15.6		0000	
4148	9130	T4 – Thyroxine	В	All	5-9	6.4-13.3	mcg/dL	999.9 or	
7170	7150	11 Inytoxiiic	ע	1 111	10-14	5.6-11.7	meg/ul	ZZ9.9	
					15+	4.5-12.0		<u> </u>	
4150	0122	E 701	Г.	. 11				999.9 or	
4150	9130	Free Thyroxine Index	В	All	All	1.4-3.8*		ZZ9.9	
		Hepatitis B Surface				Neg			
4305	9125		В	All	All	iveg			
		Antibody				3.7		1	
4315	9125	Hepatitis B Surface	В	All	All	Neg			
		Antigen							
4325	9125	Hepatitis B Core Antibody	В	All	All	Neg			
									•

	Updated June 2023						2023		
Result Id	Remark Id	Test Description	Spec Type	Gender	Age	Reference Range	Unit of Measure	Decimal Position	Reference
4335	9125	Hepatitis B Core IgM	В	All	All	Neg			
4345	9125	Hepatitis A, Total Ab	В	All	All	Neg			
4351	9125	Hepatitis A IgM	В	All	All	Neg			
4365	9125	Hepatitis C	В	All	All	Neg			
4370	9125	HCV RNA, Quant RT- PCR	В	All	All	Not Detected	Log IU/mL	9.99	
4375	9125	Hepatitis Be Antigen	В	All	All	Neg			
4385	9125	Hepatitis Be Antibody	В	All	All	Neg			
4390	9125	Hepatitis 5-1-1p/c100p Band	В	All	All	0 - 4		9 or Z	
4391	9125	Hepatitis c33c Band	В	All	All	0 – 4		9 or Z	
4392	9125	Hepatitis c22p Band	В	All	All	0 - 4		9 or Z	
4393	9125	Hepatitis NS5 Band	В	All	All	0 - 4		9 or Z	
4394	9125	Hepatitis hSOD Band	В	All	All	0 – 4		9 or Z	
4395	9125	Hepatitis RIBA-3 Interpretation	В	All	All	Pos Neg Indeter			
4922	9100	CRP – C-Reactive Protein	В	All	All	0-5	mg/L	9999.9 or ZZZZ9.9	
4924	9100	Cardiac Relative Risk	В	All	All	1.0		9.9 or Z.9	
4975	9700	DIU – Diuretic Urine Screen	U	All	All	<1500	ng/mL	Alpha X(7)	
5000	9700	BAB – Beta Blocker Urine Screen	U	All	All	<5000	ng/mL	Alpha X(7)	
5025^	9400 9700	Cocaine - Urine	U	All	All	0 = Negative 1 = Positive		9 or Z	
5050	9400 9700	Glucose – Urine	U	All	All	0.00 - 0.24	gm%	9.99 or Z.99	
5100^	9400	Marijuana – Urine	U	All	All	0 = Negative 1 = Positive		9 or Z	
5125	9700	Urine Microscopy – Granular Casts	U	All	All	0	/40 LPF	9999 or ZZZ9	
5150	9700	Urine Microscopy – Hyaline Casts	U	All	All	0-10	/40 LPF	9999 or ZZZ9	
5175	9700	Urine Microscopy – RBC	U	All	All	0-4	/HPF	9999 or ZZZ9	
5200	9700	Urine Microscopy – WBC	U	All	All	0-9	/HPF	9999 or ZZZ9	
5220	9100	Cotinine (Nicotine) - Serum	В	All	All	<25	ng/mL	Alpha X(7)	
5225	9400 9700	Cotinine (Nicotine) – Urine	U	All	All	<.50	mcg/mL	9.99 or Z.99	

							Úpdated June 2023		
Result Id	Remark Id	Test Description	Spec Type	Gender	Age	Reference Range	Unit of Measure	Decimal Position	Reference
5235	9700	HCG (Pregnancy Test) – Urine	U	All	All	Neg		Alpha X(7)	
5250	9400 9700	Protein – Urine	U	All	All	0-30	mg%	9999 or ZZZ9	
5260	9700	Albumin - Urine	U	All	All	0-3	mg/dL	9999.9 or ZZZ9.9	
5265	9700	Albumin/Creatinine Ratio	U	All	All	0-30		9999.9 or ZZZ9.9	
5275	9700	Specific Gravity	U	All	All	1.003-1.035		9.999	
5300^	9400	Toxicology – Amphetamine	U	All	All	0 = Negative 1 = Positive		9 or Z	
5350^	9400	Toxicology – Codeine	U	All	All	0 = Negative 1 = Positive		9 or Z	
5450^	9400	Toxicology – Morphine	U	All	All	0 = Negative 1 = Positive		9 or Z	
5475^	9400	Toxicology – Oxycodone	U	All	All	0 = Negative 1 = Positive		9 or Z	
5480^	9400	Toxicology - Oxymorphone	U	All	All	0 = Negative 1 = Positive		9 or Z	
5525^	9400	Toxicology – Phencyclidine	U	All	All	0 = Negative 1 = Positive		9 or Z	
5575^	9400	Toxicology – Methamphetamine	U	All	All	0 = Negative $1 = $ Positive		9 or Z	
5590^	9400	Toxicology – Codeine	U	All	All	0 = Negative 1 = Positive		9 or Z	
5595^	9400	Toxicology – Hydrocodone	U	All	All	0 = Negative 1 = Positive		9 or Z	
5600^	9400	Toxicology – Morphine	U	All	All	0 = Negative 1 = Positive		9 or Z	
5605^	9400	Toxicology – Hydromorphone	U	All	All	0 = Negative 1 = Positive		9 or Z	
5660^	9400	Toxicology – Fentanyl	U	All	All	0 = Negative $1 = $ Positive		9 or Z	
5662^	9400	Toxicology – Norfentanyl	U	All	All	0 = Negative $1 = $ Positive		9 or Z	
6000	9800	Urine Antibody Screen Interpretation	U	All	All	Neg		Alpha X(7)	
6005		Urine Temperature	U	All	All	90.5-99.6	°F	999.9 or Z99.9	
6007	9800	Urine Antibody Screen Interpretation (U.S.)	U	All	All	Neg			
6015	9400 9700	Urine Adulterant – Creatinine	U	All	All	27.0-260.0 Adulterant Range: >5.0	mg/dL	9999.9 or ZZZ9.9	
6020	9400 9700	Urine Adulterant – pH	U	All	All	5.0 – 8.0		99.9 or Z9.9	
6044	9175	Dried Blood – GGTP	D	All	All	< 65	U/L	9999 or ZZZ9	
6055^	9175	Dried Blood – Cocaine	D	All	All	0 = Negative 1 = Positive		9,Z	
6060	9180	Dried Blood – A1C	D	All	All	3.0-6.0	%	99.9 or Z9.9	
6520	9875	Oral Fluid Antibody Screen Interpretation	S	All	All	Neg		Alpha X(7)	
6525	9875	Oral Fluid Confirmation	S	All	All	Y		Alpha X(7)	

							ι	pdated June	2023
Result Id	Remark Id	Test Description	Spec Type	Gender	Age	Reference Range	Unit of Measure	Decimal Position	Reference
6550	9880	Oral Fluid Cocaine	S	All	All	10/SCRN 10/CONF	ng/mL	Alpha X(7)	
6555	9880	Oral Fluid Cotinine	S	All	All	<10	ng/mL	Alpha X(7)	
6560	9875	Oral Fluid Antibody Screen Interpretation (U.S.)	S	All	All	Neg			
6600		Applicant Weight		All	All		lb	999.9 or ZZ9.9	Exam Data
6605		Applicant Height		All	All		feet	99 or Z9	Exam Data
6610		Applicant Height		All	All		inches	99.9 or Z9.9	Exam Data
6615		Blood Pressure – Systolic		All	All			999, ZZ9 or ZZZ	Exam Data
6620		Blood Pressure – Systolic 2		All	All			999, ZZ9 or ZZZ	Exam Data
6625		Blood Press – Systolic 3		All	All			999, ZZ9 or ZZZ	
6630		Blood Pressure – Diastolic		All	All			999, ZZ9 or ZZZ	Exam Data
6635		Blood Pressure – Diastolic 2		All	All			999, ZZ9 or ZZZ	Exam Data
6640		Blood Pressure – Diastolic 3		All	All			999, ZZ9 or ZZZ	Exam Data
6645		Pulse Rate – Standard – 1		All	All			999, ZZ9 or ZZZ	Exam Data
6650		Pulse Rate – Standard – 2		All	All			999, ZZ9 or ZZZ	Exam Data
6655		Pulse Rate – Standard – 3		All	All			999, ZZ9 or ZZZ	Exam Data
6660		Pulse Rate – Irregular – 1		All	All			999, ZZ9 or ZZZ	Exam Data
6665		Pulse Rate – Irregular – 2		All	All			999, ZZ9 or ZZZ	Exam Data
6670		Pulse Rate – Irregular – 3		All	All			999, ZZ9 or ZZZ	Exam Data
6672		BMI – Body Mass Index		All	All	18.5 – 24.9		9.9 or Z9.9	Exam Data
6676	9100	CKDEPI-Creat GFR	В	All	All	>=60	mL/min	999 or ZZ9	Effective 6/1/2023
6677	9100	CKDEPI Creat EGFR AFR	₽	All	All	>=60	mL/min	999 or ZZ9	Phasing out effective 6/1/2023
6678	9100	CKD EPI Cystatin C EGFR	В	All	All	>=60	mL/min	999 or ZZ9	
6679		Body Surface Area		All	All			9.99 or Z.99	Effective 6/1/2023
7120	9400 9700	Protein/Creatinine Ratio	U	All	All	0-0.20		999.99 or ZZ9.99	
7125	9400 9700	Leukocyte Esterase Screen	U	All	All	Neg		ZZZZ9	
7130	9400 9700	Hemoglobin Screen	U	All	All	Neg		ZZZZ9	
8105	9890	CDT – Quantitative		F M	All	0 - 66 0 - 35	mcg/mL	ZZ9.9Z or 999.99	
8901		Tobacco Use		All	All	Yes/No			Tobacco

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Result Id	Remark Id	Test Description	Spec Type	Gender	Age	Reference Range	Unit of Measure	Decimal Position	Reference
8902		Tobacco Product Use – Cigarette		All	All	Yes/No			Tobacco
8903		History of Hypertension		All	All	Yes/No			Part II
8904		History of Diabetes		All	All	Yes/No			Part II
8905		History of Proteinuria/Albuminuria		All	All	Yes/No			Part II
8906		Medication Use		All	All	Yes/No			Medication
8907		Tobacco Product Use – Cigar		All	All	Yes/No			Tobacco
8908		Tobacco Product Use – Pipe		All	All	Yes/No			Tobacco
8909		Tobacco Product Use – Smokeless		All	All	Yes/No			Tobacco
8910		Nicotine – Delivery System		All	All	Yes/No			Tobacco
8911		Cardiac History		All	All	Yes/No			Part II

[^]indicates a sensitive result

Appendix C

- ➤ Units of Measure
- ➤ Results Exceeding Reporting Limit
- > Remark Ids and Associated Result Ids
- ➤ Replacement Remark Descriptions
- ➤ Interpretation of Remarks
- > Ratio Calculations
- ➤ Remarks to Associate With Results
- ➤ Results Not Reported If Not Performed
- > Serum Appearances
- > Exam Questions

Units of Measure

UNIT	DESCRIPTION			
%	percent			
μ^3	cubic microns			
°F	degrees Fahrenheit			
μIU/mL	micro international units per milliliter			
μmol/L	micromoles per liter			
/40 lpf	per 40 low powered field			
/hpf	per high powered field			
g/dL	grams per deciliter			
gm%	grams percent			
Log IU/mL	Log international units per milliliter			
mcg/dL	micrograms per deciliter			
mcg/mL	micrograms per milliliter			
meq/L	milliequivalents per liter			
mg%	milligrams percent			
mg/dL	milligrams per deciliter			
mg/L	milligrams per liter			
mL/min	milliliter per minute			
mmol/L	millimoles per liter			
pg	picograms			
U/L	international units per liter			

Results Exceeding Reporting Limit

Reporting a value for some results beyond a specific limit may have limited (if any) clinical significance. In the following cases, a specific result value is reported which is used to designate that the normal reporting limit has been exceeded.

Result Id	Minimum/Maximum Interpreted	CPU to CPU [®] Raw Data Displayed	Lab <i>One</i> Text Reported Value
1275	2	0002000	<3
1300	0	0000000	<0.1
1325	1	0001000	<2
1325	201	0201000	>200
1330	35	0035000	<36
1330	9999	9999000	>9999
1525	9	0009000	<10
1525	9999	9999000	>9999
1600	4	0004000	<5
1600	201	0201000	>200
2100	3.9	0003900	<4.0
2100	14.0	0014100	>14.0
2680	0.24	0000240	< 0.25
2680	96.1	0096100	>96.0
2685	0.03	0000030	< 0.04
2685	100.01	0100010	>100.00
2705	301	0301000	>300
3095	0.48	0000480	< 0.49
3095	13.81	0013810	>13.80
3350	149	0149000	<150
3350	301	0301000	>300
3693	0	0000000	<0.1
3693	12.1	0012100	>12.0
3701	24.0	0024000	<25.0
3701	800.1	0800100	>800.0
3703	0	0000000	< 0.01
3950	1.2	0001200	<1.3
3952	1.9	0001900	<2.0

Result Id	Minimum/Maximum Interpreted	CPU to CPU [®] Raw Data Displayed	Lab <i>One</i> Text Reported Value
4144	14.9	0014900	<15.0
4144	50.1	0050100	>50.0
4145	0.4	0000400	< 0.5
4145	20.1	0020100	>20.0
4148	0.6	0000600	< 0.7
4148	40.1	0040100	>40.0
4370	0.00	0000000	<1.18 Not Detected
4370	1.17	0001170	<1.18 Detected
4370	8.00	0008000	>7.99
4922	0.2	0000200	<0.3
4922	10.1	0010100	>10.0
5050	0.00	0000000	< 0.01
5050	1.1	0001100	>1.0
5225	0.09	0000090	< 0.10
5225	1.01	0001010	>1.00
5260	0.0	0000000	<0.1
5260	600.1	0600100	>600.0
5275	1.036	0001036	>1.035
6020	3.9	0003900	<4.0
6020	9.1	0009100	>9.0
6044	59	0059000	<60
6044	361	0361000	>360
6050	99	0099000	<100
6050	301	0301000	>300
6060	2.9	0002900	<3.0
6060	18.1	0018100	>18.0

Remark Ids and Associated Result Ids

Remark Id	Description	Associated Result Ids		
		1050		
9000		1075		
		1100		
	CDC	1125		
	CBC	1150		
		1175		
		1200		
		1225		
9010	Chain of Custody Comments			
9020	Client Quoteback			
	1250			
		1251		
		1275		
		1300		
		1325		
		1330		
		1350		
		1375		
		1380		
		1400		
		1425		
		1450		
		1475		
	Blood Chemistry	1500		
0100		1525		
9100		1560		
		1575		
		1600		
		1625		
		1630		
		1635		
		2100		
		3095		
		3664		
		4922		
		4924		
		5220		
		6676		
		6677		
		6678		
		4305		
9125		4303		
		4313		
		4325		
	Honotitic	4345		
	Hepatitis	4345		
		4365		
		4370		
		4375		

Remark Id	Description	Associated Result Ids
	•	4385
		4390
		4391
		4392
		4393
		4394
		4395
		3693
		3701
9130	Thyroid	3703
7130	Thyroid	4144
		4148
		4150
		3340
9175	Dried Blood Profile - Chemistry	3350
	,	6044
		6055
9180	Dried Blood A1C	6060
9200	Serum HIV – Antibody Screen	2125
9200	Scrum III v – Antibody Screen	2600
9250	Medications	
9275	Examiner Comments	
**9300	Serum Appearance	
9375	Dried Blood Profile – HIV	3135
		3289
		4010
		4012
		4014
		5025
		5050
		5100
		5225
		5250
		5300
		5350
		5450
9400	Toxicology	5475
		5480
		5525
		5575
	5590 5595 5600	
		5605
		5660
		5662
		6015
		6020
		7120

Remark Id	Description	Associated Result Ids
		7125
		7130
9500	Applicant Message	
		1650
		1675
		1700
		1725
		1750
9650	Differential	1775
9030	Differential	1800
		1825
		1850
		1875
		1900
		4975
		5000
		5025
		5050
		5100
		5125
		5150
		5175
		5200
9700	Urine	5225
9700	Office	5235
		5250
		5260
		5265
		5275
		6015
		6020
		7120
		7125
		7130
		2685
9750	PSA/Tumor Marker	3950
9/30	r SA/ I UIIIOF IVIATKET	3952
		3959
0000	Hala HINA (1 1 C	6000
9800	Urine HIV Antibody Screen	6007
*9850	Urine Adulterant	
9875	Oral Fluid HIV Antibody Screen	6520 6525
	-	
9880	Oral Fluid Drug	6550
		6555
0000	A111 N.C. 1	2705
9890	Alcohol Marker	2720
	CD 1 C 1	2722

^{*}Refer to the "Interpretation of Remarks: section for more details.

** Refer to the "Serum Apperance Values: section for more details.

Replacement Remark Descriptions

The following are possible replacement remarks for the result fields of Record Types 010-099. Each test result may have any of these replacement remarks.

REMARK	DESCRIPTION		
DODDI NE	BORDERLINE		
BORDLNE CBLT	CDT RESULTS CONFIRMED		
CONS	QNS TO CONFIRM		
EXAL	NO ORAL FLUID TESTS PERFORMED – KIT EXPIRED		
EXHI	NO ORAL FLUID HIV TEST PERFORMED – KIT EXPIRED		
FEW	FEW		
INDETER	INDETERMINATE		
MAN	MANY		
MAR	MARKED		
MOD	MODERATE		
MOHE	MODERATE HEMOLYSIS		
MOLI	MODERATE LIPEMIA		
N	NO. CLASS G IMMUNOGLOBULIN LEVEL BELOW VALID		
	LIMITS[Oral Fluid Specimen Validity]		
NCAL	NOT CALCULATED		
NEG	NEGATIVE		
NG	NOT GIVEN		
NI	INVALID DUE TO ICTERUS		
NO	NO		
NONE	NONE		
NOR	NORMAL		
NP	NOT PERFORMED		
NSR	NO SPECIMEN RECEIVED		
NT	TEST NOT PERFORMED-NO SPECIMEN		
NVG	RESULT NOT VALID DUE TO GLYCOLYSIS		
NVH	RESULT NOT VALID DUE TO LIBEMIA		
NVL OCLT	RESULT NOT VALID DUE TO LIPEMIA OCCULT BLOOD DETECTED		
OTR	OUT OF RANGE		
OUT	OUTSIDE REPORTABLE LI MITS		
POS	POSITIVE OR REACTIVE		
QNS	QUANTITY NOT SUFFICIENT		
REF	REFER TO REMARK BELOW		
SEHE	SEVERE HEMOLYSIS		
SELI	SEVERE LIPEMIA		
SIBT	UNABLE TO PERFORM URINE HIV-COLLECTOR USED IMPROPER		
	NON HIV CONTAINER		
SLHE	SLIGHT HEMOLYSIS		
SLLI	SLIGHT LIPEMIA		
SPCE	SPECIMEN NOT CENTRIFUGED		
SPPO	SPECIMEN NOT POURED OFF		
SPRM	SPERM		
SRC	SMALL ROUND CELL		
SUFA	SPECIMEN UNSUITABLE FOR ANALYSIS		

REMARK	DESCRIPTION
TNSA	T-CELL NOT SUITABLE FOR ANALYSIS
TNTC	TOO NUMEROUS TO COUNT
TRIG	INVALID WHEN TRIGLYCERIDES >400
XTAL	CRYSTAL
Y	YES. CLASS G IMMUNOGLOBULIN LEVEL WITHIN VALID LIMITS. SPECIMEN VALID[Oral Fluid Specimen Validity]
YES	YES
YLC	YEAST

Interpretation of Remarks

❖ For Remark Id 9200, when the value reported is MSG01, the following statement should be reported by your application:

T-CELL ABNORMALITIES CAN BE CAUSED BY A VARIETY OF FACTORS INCLUDING, BUT NOT LIMITED TO: VIRAL INFECTIONS, EXPOSURE TO CHEMICAL TOXINS, ANITMICROBIAL TREATMENTS, AIDS, BONE MARROW TRANSPLANTS.

- ❖ Any negative urine adulterant result will report in Remark Id 9850 with the following statement: ADULTERANT TESTS WITHIN NORMAL RANGE
- ❖ For all specimens collected in New York, the following should be associated:

THIS INFORMATION HAS BEEN DISCLOSED TO YOU FROM CONFIDENTIAL RECORDS WHICH ARE PROTECTED BY STATE LAW. STATE LAW PROHIBITS YOU FROM MAKING ANY FURTHER DISCLOSURE OF THIS INFORMATION WITHOUT THE SPECIFIC WRITTEN CONSENT OF THE PERSON TO WHOM IT PERTAINS, OR AS OTHERWISE PERMITTED BY LAW.

New York state law mandates the following comment be reported on all Urine HIV results where the urine was collected in a non-regulated container:

COLLECTED IN A URINE VIAL FROM A BLOOD COLLECTION KIT. FOR INSURANCE RISK ASSESSMENT ONLY – NOT FOR DIAGNOSTIC PURPOSES.

Ratio Calculations

While most of the results (including ratios) in the CPU-to-CPU® system are determined and obtained directly from the equipment used to perform the tests, a few result values are calculated using result values for other related Result Ids. The following is a list of calculated ratios and the results used in computing them.

RESULT ID	CALCULATION
1030	Result Id 1000 divided by Result Id 1025
1625	Result Id 1350 divided by Result Id 1600
1635	Result Id 1630 divided by Result Id 1600
2375	Result Id 2225 divided by Result Id 2250
7120	Result Id 5250 divided by Result Id 6015
5265	(Result Id 5260 divided by Result Id 6015)*1000

Note: Calculations are usually performed using the related result values before they are rounded or truncated and may be computed using a greater number of significant digits than reported. Using the reported values in the related results to recreate the calculations may not result in values that will equate to those supplied in the Result Ids for the ratios.

Remarks to Associate with Results

For Free PSA, Result Id 3959, the following statement should be displayed:

< 10	% FREE PSA	INCREASED RISK OF CANCER
10 - 25	% FREE PSA	INTERMEDIATE RISK OF CANCER
>25	% FREE PSA	DECREASED RISK OF CANCER

THE PERCENT FREE PSA IS INVERSELY PROPORTIONAL TO THE RISK OF PROSTATE CANCER. PERCENT FREE PSA IS NOT ABSOLUTE EVIDENCE OF MALIGNANCY. PERCENT FREE PSA IS PERFORMED USING THE BECKMAN COULTER ACCESS II. VALUES OBTAINED WITH DIFFERENT ASSAY METHODS MAY NOT BE INTERCHANGEABLE.

For CEA (Carcinoembryonic Antigen) Result Id 3952, the following statement should be displayed:

CEA IS NOT ABSOLUTE EVIDENCE OF MALIGNANCY. CEA PERFORMED USING BAYER DIAGNOSTICS CENTAUR CHEMILUMINSCENCE ASSAY. VALUES OBTAINED WITH DIFFERENT ASSAY METHODS MAY NOT BE INTERCHANGEABLE.

For AFP (Alpha-Fetoprotein) Result Id 3950, the following statement should be displayed:

AFP PERFORMED USING THE BECKMAN COULTER CHEMILUMINESCENT METHOD. VALUES OBTAINED FROM DIFFERENT ASSAY METHODS CANNOT BE USED INTERCHANGEABLY. AFP LEVELS, REGARDLESS OF VALUE, SHOULD NOT BE INTERPRETED AS ABSOLUTE EVIDENCE OF THE PRESENCE OF ABSENCE OF DISEASE.

For Urine HIV Result ID 6007, the following statement should be displayed:

THE URINE HIV-1 TEST MAY MISS 1 TO 2% OF ALL TRUE POSITIVES. THE TEST MAY GENERATE UP TO 6% FALSE POSITIVES WHEN TESTING SUBJECTS AT HIGH RISK FOR HIV-1 INFECTION AND 10% FALSE POSITIVES FOR INDIVIDUALS WITH OTHER MEDICAL CONDITIONS.

For HCV RNA Quant RT-PCR, Result ID 4370, the following statement should be displayed:

THIS TEST WAS PERFORMED USING REAL-TIME POLYMERASE CHAIN REACTION.

THE ANALYTICAL PERFORMANCE CHARACTERISTICS OF THIS ASSAY HAVE BEEN DETERMINED BY QUEST DIAGNOSTICS. THE MODIFICATIONS HAVE NOT BEEN CLEARED OR APPROVED BY THE FDA. THIS ASSAY HAS BEEN VALIDATED PURSUANT TO THE CLIA REGULATIONS.

Results Not Reported if Not Performed

While some Result Ids may be reported along with a result value of "NP" to indicate that the test was NOT PERFORMED, the following Result Ids will NOT be reported if the test was not performed:

3135	5275	6050	6620	6645	6670
5125	6005	6600	6625	6650	6672
5150	6015	6605	6630	6655	
5175	6020	6610	6635	6660	
5200	6044	6615	6640	6665	

Serum Appearance Values

The following are examples of serum appearance values that will be sent in Remark 9300 of Record Type 100-199.

MODERATE HEMOLYSIS
MODERATE LIPEMIA
NONE
NORMAL
SEVERE HEMOLYSIS
SEVERE LIPEMIA
SLIGHT HEMOLYSIS
SLIGHT LIPEMIA
SPECIMEN NOT CENTRIFUGED
SPECIMEN NOT POURED OFF

Exam Questions

Exam questions are collected by the Paramed/Examiner, recorded on the Insurance Id Slip, submitted to Exam*One*, data entered, and then reported to the client using their corresponding Result Ids. They are not results generated from testing performed by Exam*One*. There is an extra fee associated with capturing and reporting exam questions, therefore each service will be added only upon request by your Medical Underwriting Department.

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Appendix D

➤ Agreement to Provide Sensitive Test Results via Electronic Transmission



Agreement to Provide Sensitive Test Results via Electronic Transmission

This agreement will serve to provide Exam*One* authorization to electronically transmit sensitive test results to your company. Sensitive test results may include results containing positive HIV, positive cocaine and/or other positive drugs of abuse on your insurance applicants. This agreement will cover the transmission of results to an Exam*One* System (software and/or hardware) or CPU-to-CPU® (direct transmission or via a third party network).

Exam*One* recommends that Company has the following security measures in place to restrict access to these sensitive test results to ensure the confidentiality of this information:

- 1. Restrict access to the physical hardware to those individuals needing access to these results and who are authorized to view applicants' results, if possible.
- 2. Utilize the password protection provided by the Exam*One* Portal. This password should be changed periodically, in accordance with recommended security best practices. For CPU-to-CPU[®] clients, we suggest you construct your software application to enforce password protection that would prevent unauthorized individuals from gaining access to this data
- 3. Control the knowledge of the password. Maintain a list of those authorized to know this information and review the list periodically.

ExamOne Portal

4. Initiate procedures that will protect printed copies of the results from being available/reviewed by unauthorized persons.

Cocaine/Drugs of Abuse	Cocaine/Drugs of Abuse	
HIV	HIV	
By signing below, the Company requests and au manner described above, and Company assumes results. The individual signing represents that he	s the responsibility for the confidentiality and s	security of those
Accepted and agreed to on this	s day of 20	-
Signature:		_
Name:		_
Title:		_
Company:		

CPU-to-CPU® (Includes ANSI 186)