HL7 v2.5 ORU^R01^ORU_R01 Message: Incorporation of Laboratory Results						
Test Case ID	LRI_1.2_1.1-GU					
Juror ID						
Juror Name						
HIT System Tested						
Inspection Date/Time						
Inspection Settlement (Pass/Fail)	Pass	Fail				
Inspection Settlement (1 ass/1 an)						
Reason Failed						
Juror Comments						

This Test Case-specific Juror Document provides a checklist for the Tester to use during testing for assessing the Health IT Module's ability to display and incorporate required data elements from the information received in the LRI message. Additional data from the message or from the Health IT Module are permitted to be displayed and incorporated by the Module. Grayed-out fields in the Juror Document indicate where no data for that data element were included in the LRI message for the given Test Case.

The format of the Display Verification section of this Juror Document is for ease-of-use by the Tester and does not indicate how the Health IT Module display must be designed.

### **Display Verification**

#### **Legend for Display Requirement**

Data in **bold red** text: HIT Module must display exact version of stored data

Data in bold black italics text: HIT Module must display exact version of data received in the LRI message

Data in regular text: HIT Module may display equivalent version of stored data

Patient Information - Display Verification								
Patient Identifier Patient Name DOB Sex Race Tester Comment								
PATID1236 Janet Anderson 09/30/1986 F White								
When a given patient has more than one Pa for the context (e.g., inpatient ID Number		nay display the ID Number	er that is mo	ost appropriate				

	Lab Results - Display Verification								
Test Performed:	est Performed: Erythrocyte sedimentation rate								
Test Report Date:	09/26/2015	14:05:	51						
Result Report Status	X								
Result Observation Name	Result Value								
Erythrocyte sedimentation rate	Test could not be performed, see Note for details				X	09/25/2015 14:00:		09/26/2015 13:05:50	
Note	Blood in tube was clotted, resulting in a rejection of the specimen and leaving the lab unable to perform this test. Please resubmit a new specimen, if test is still desired.								

Performing Organization Information - Display Verification						
Data Element Name	Data Element Name Data					
Organization Name	Century Hospital					
Organization Address						
Street address	2070 Test Park					
Other designation						
City	Los Angeles					
State	CA					
Zip code	90067					

Performing Organization Medical Director Information - Display Verification									
Data Element Name	Tester Comment								
Medical Director Name	Medical Director Name								
Family Name									
Surname	Knowsalot								
Given Name	Phil								
Second and Further Given Names or Initials Thereof	J.								
Suffix (e.g., JR or III)	III								
Prefix (e.g., DR)	Dr.								

Specimen Information - Display Verification								
Data Element Name	Tester Comment							
Specimen Type(Specimen Source)	Blood Specimen							
Specimen Collection Date/Time - Start	09/25/2015 14:00:							
Specimen Collection Date/Time - End								
Specimen Reject Reason	Blood specimen clotted							
Specimen Condition	blood specimen clotted							

Order Information - Display Verification							
Data Element Name	Data Element Name Data						
Relevant Clinical Information							
Placer Order Number Entity ID	ORD723222-1						
Ordering Provider							
Family Name							
Surname	Radon						
Given Name	Nicholas						
Second and Further Given Names or Initials Thereof							
Suffix (e.g., JR or III)							
Prefix (e.g., DR)							

# **Incorporate Verification**

### Legend for Store Requirement

S-EX: Store exact

 $S-TR-R: Translate \ and \ store \ translation \ (exact \ value \ can \ be \ re-created \ from \ translation \ any \ time)$ 

S-EX-A: Store exact by association

S-RC: Process and re-create

S-EQ : Store equivalent

(See "Instructions to Testers for Verification of Store Requirements" at the end of this Juror Document for additional details.)

	Patient Information Details- Incorporate Verification						
Location	Data Element Name	Store Requirement	Data	Tester Comment			
PID-3	Patient Identifier List						
PID-3.1	ID Number	S-EX-A	PATID1236				
PID-3.4	Assigning Property						
PID-3.4.1	Namespace ID	S-EX-A					
PID-3.4.2	Universal ID	S-EX-A	2.16.840.1.113883.3.72.5.30.2				
PID-3.4.3	Universal ID Type	S-EX-A	ISO				
PID-3.5	Identifier Type Code	S-RC	MR				
PID-5	Patient Name						
PID-5.1	Family Name						
PID-5.1.1	Surname	S-EX-A	Anderson				
PID-5.2	Given Name	S-EX-A	Janet				
PID-5.3	Second and Further Given Names or Initials Thereof	S-EX-A					
PID-5.4	Suffix (e.g., JR or III)	S-EX-A					
PID-5.7	Name Type Code	S-RC	L				
PID-7	Date/Time of Birth						
PID-7.1	Time	S-EQ	09/30/1986				
PID-8	Administrative Sex	S-TR-R	F				
PID-10	Race	_					
PID-10.1	Identifier	S-RC	2106-3				
PID-10.2	Text	S-RC	White				
PID-10.3	Name of Coding System	S-RC	HL70005				

		Order Informa	ation - Incorporate Verifica	ation
Location	Data Element Name	Store Requirement	Data	Tester Comment
	Placer Order Number			
ORC-2.1/OBR- 2.1	Entity Identifier	S-EX-A	ORD723222-1	
ORC-2.2/OBR- 2.2	Namespace ID	S-EX-A		
ORC-2.3/OBR- 2.3	Universal ID	S-EX-A	2.16.840.1.113883.3.72.5.24	
ORC-2.4/OBR- 2.4	Universal ID Type	S-EX-A	ISO	
ORC-3/OBR-3	Filler Order Number			
ORC-3.1/OBR- 3.1	Entity Identifier	S-EX	R-783274-1	
ORC-3.2/OBR- 3.2	Namespace ID	S-EX-A		
ORC-3.3/OBR- 3.3	Universal ID	S-EX-A	2.16.840.1.113883.3.72.5.25	
ORC-3.4/OBR- 3.4	Universal ID Type	S-EX-A	ISO	
ORC-12/OBR-16	Ordering Provider			
ORC-12.1/OBR- 16.1	ID Number	S-RC	5742200012	
ORC- 12.2/OBR-16.2	Family Name			
ORC- 12.2.1/OBR-16.2.1	Surname	S-RC	Radon	
16.3	Given Name		Nicholas	
10.4	Second and Further Given Names or Initials Thereof	S-RC		
	, , ,	S-RC		
ORC-12.6/OBR- 16.6	Prefix (e.g., DR)	S-RC		
ORC- 12.9/OBR-16.9	Assigning Authority			
ORC- 12.9.1/OBR-16.9.1	Namespace ID	S-EX-A		
ORC- 12.9.2/OBR-16.9.2	Universal ID	S-EX-A	2.16.840.1.113883.4.6	
ORC- 12.9.3/OBR-16.9.3	Universal ID Type	S-EX-A	ISO	
ORC- 12.10/OBR-16.10	Name Type Code	S-RC	L	
ORC- 12.13/OBR-16.13	Identifier Type Code	S-RC	NPI	

		Store		
Location	Data Element Name	Store Requirement	Data	Tester Comment
OBX-23	Performing Organization Name			
OBX-23.1	Organization Name (Note 1)	S-TR-R	Century Hospital	
OBX-23.6	Assigning Authority (Note 2)			
OBX-23.6.1	Namespace ID	S-EX-A	CLIA	
OBX-23.6.2	Universal ID	S-EX-A	2.16.840.1.113883.4.7	
OBX-23.6.3	Universal ID Type	S-EX-A	ISO	
OBX-23.7	Identifier Type Code	S-RC	XX	
OBX-23.10	Organization Identifier	S-TR-R	24D9871327	
OBX-24	Performing Organization Address			
OBX-24.1	Street Address			
OBX-24.1.1	Street or Mailing Address	S-EX-A	2070 Test Park	
OBX-24.2	Other Designation	S-EX-A		
OBX.24.3	City	S-EX-A	Los Angeles	
OBX-24.4	State or Province	S-EX-A	CA	
OBX-24.5	Zip or Postal Code	S-EX-A	90067	
OBX-24.6	Country	S-TR-R	USA	
OBX-25	Performing Organization Medical Director			
OBX-25.1	ID Number	S-RC	5432178916	
OBX-25.2	Family Name			
OBX-25.2.1	Surname	S-TR-R	Knowsalot	
OBX-25.3	Given Name	S-TR-R	Phil	
OBX-25.4	Second and Further Given Names or Initials Thereof	S-TR-R	J.	
OBX-25.5	Suffix (e.g., JR or III)	S-TR-R	III	
OBX-25.6	Prefix (e.g., DR)	S-TR-R		
OBX-25.9	Assigning Authority (Note 2)			
OBX-25.9.1	Namespace ID	S-EX-A	NPI	
OBX-25.9.2	Universal ID	S-EX-A	2.16.840.1.113883.4.6	
OBX-25.9.3	Universal ID Type	S-EX-A	ISO	
OBX-25.10	Name Type Code	S-RC	L	
OBX-25.13	Identifier Type Code	S-RC	NPI	

Note 1 - The HIT Module must store the Organization Name or be able to recreate it. If the HIT Module is able to demonstrate Organization Name: ID is always 1:1, then the HIT Module is permitted to store and recreate (S-TR-R).

Note 2 - Determine requirement for support of 2nd component or 3rd and 4th component based on the EI or HD Profile

	Order Information (cont'd) - Incorporate Verification					
Location	Data Element Name	Store Requirement	Data	Tester Comment		
OBR-4	Universal Service Identifier (Note 1)					
OBR-4.1	Identifier	S-TR-R	30341-2			
OBR-4.2	Text	S-EX-A	Erythrocyte sedimentation rate			
OBR-4.3	Name of the Coding System	S-RC	LN			
OBR-4.4	Alternate Identifier	S-TR-R	500			
OBR-4.5	Alternate Text	S-EX-A	Erythrocyte sedimentation rate			
OBR-4.6	Name of Alternate Coding System	S-RC	99USL			
OBR-4.9	Original Text	S-EX	Erythrocyte sedimentation rate			
OBR-7/SPM-17.1	Observation Date/Time					
OBR-7.1/SPM- 17.1.1	Time	S-EQ	09/25/2015 14:00:			
OBR-8/SPM-17.2	Observation End Date/Time					
OBR-8.1/SPM- 17.2.1	Time	S-EQ				
OBR-13	Relevant Clinical Information					
OBR-13.1	Identifier	S-TR-R				
OBR-13.2	Text	S-EX-A				
OBR-13.3	Name of the Coding System	S-RC				
OBR-13.9	Original Text	S-EX				
OBR-22	Results Rpt/Status Chng - Date/Time					
OBR-22.1	Time	S-EQ	09/26/2015 14:05:51			
OBR-25	Result Status	S-TR-R	X			

Note 1 -Store the <u>Identifier</u> and the <u>Text</u> for each populated triplet using the S-EX-A, S-TR-R, or S-EX store requirement as indicated. If <u>Original Text</u> field is populated, MUST store the exact data received.

	Result Information - Incorporate Verification					
Location	Data Element Name	Store Requirement	Data	Tester Comment		
OBX-3	Observation Identifier (Note 1)					
OBX-3.1	Identifier	S-TR-R	30341-2			
OBX-3.2	Text	S-EX-A	Erythrocyte sedimentation rate			
OBX-3.3	Name of the Coding System	S-RC	LN			
OBX-3.4	Alternate Identifier	S-TR-R	815117			
OBX-3.5	Alternate Text	S-EX-A	ESR			
OBX-3.6	Name of Alternate Coding System	S-RC	99USL			
OBX-3.9	Original Text	S-EX	Erythrocyte sedimentation rate			
OBX-5	Observation Value	S-EX	Test could not be performed, see Note for details			
OBX-6	Units (Note 2)					
OBX-6.1	Identifier	S-TR-R				
OBX-6.2	Text	S-TR-R				
OBX-6.3	Name of the Coding System	S-RC				
OBX-6.4	Alternate Identifier	S-TR-R				
OBX-6.5	Alternate Text	S-TR-R				
OBX-6.6	Name of Alternate Coding System	S-RC				
OBX-6.9	Original Text	S-EX				
OBX-7	Reference Range	S-EX				
OBX-8	Abnormal Flags	S-TR-R				
OBX-11	Observation Result Status	S-TR-R	Х			
OBX-14	Date/Time of the Observation					
OBX-14.1	Time	S-EQ	09/25/2015 14:00:			
OBX-19	Date/Time of the Analysis					
OBX-19.1	Time	S-EQ	09/26/2015 13:05:50			

Note 1 - Store the <u>Identifier</u> and the <u>Text</u> for each populated triplet using the S-EX-A, S-TR-R, or S-EX store requirement as indicated. If <u>Original Text</u> field is populated, MUST store the exact data received.

Note 2 - If both UOM triplets are populated, receiver may choose to store the data received in either triplet; translations must result in equivalent UOM that do not require a change in the numeric result.

Note - Incorporate Verification							
Location	Data Element Name	Store Requirement	Data	Tester Comment			
NTE-3	Note	S-EX	Blood in tube was clotted, resulting in a rejection of the specimen and leaving the lab unable to perform this test. Please resubmit a new specimen, if test is still desired.				

Specimen Information - Incorporate Verification						
Location	Data Element Name	Store Requirement	Data	Tester Comment		
SPM-4	Specimen Type (Note 1)					
SPM-4.1	Identifier	S-TR-R	119297000			
SPM-4.2	Text	S-EX-A	BLD			
	Name of the Coding System	S-RC	SCT			
SPM-4.4	Alternate Identifier	S-TR-R	BldSpc			
SPM-4.5	Alternate Text	S-EX-A	Blood			
	Name of Alternate Coding System	S-RC	99USL			
SPM-4.9	Original Text	S-EX	Blood Specimen			
	Specimen Reject Reason (Note 1)					
SPM-21.1	Identifier	S-TR-R	RC			
SPM-21.2	Text	S-EX-A	Clotting			
	Name of the Coding System	S-RC	HL70490			
SPM-21.4	Alternate Identifier	S-TR-R	С			
SPM-21.5	Alternate Text	S-EX-A	Clotting			
	Name of Alternate Coding System	S-RC	99USL			
SPM-21.9	Original Text	S-EX	Blood specimen clotted			
	Specimen Condition (Note 1)					
SPM-24.1	Identifier	S-TR-R	CLOT			
SPM-24.2	Text	S-EX-A	Clotted			
	Name of the Coding System	S-RC	HL70493			
SPM-24.4	Alternate Identifier	S-TR-R	С			
SPM-24.5	Alternate Text	S-EX-A	Clotted			
	Name of Alternate Coding System	S-RC	99USL			
SPM-24.9	Original Text	S-EX	blood specimen clotted			

Note 1 - The HIT must store the <u>Identifier</u> and the <u>Text</u> for each populated triplet using the S-EX-A, S-TR-R, or S-EX store requirement as indicated. If <u>Original Text</u> field is populated, MUST store the exact data received.

## **Instructions to Testers for Verification of Store Requirements**

Note: The HIT Module being tested is always allowed to incorporate/store the exact data received in the LRI message even if a given Store Requirement does not explicitly state that the HIT Module is permitted to do so.

Store Requirement	Definition	Instructions for Verification of Requirement During Conformance Testing	
S-EX	Store Exact	The HIT Module being tested must be designed to incorporate/store only the exact data received in the LRI message.  • Tester must verify that the HIT Module being tested incorporates/stores in the patient's laboratory result record only the exact data received in the LRI message, and that the HIT Module does not just store an equivalent of that exact data or just a pointer to the exact data.	
S-EX-A	Store exact by association	The HIT Module being tested must be designed (1) to incorporate/store the exact data received in the LRI message OR (2) to use a pointer to a location (e.g., file/table in or accessible to the HIT Module) where the exact data can be obtained.  • Tester must verify that the HIT Module being tested incorporates/stores in the patient's laboratory result record the exact data received in the LRI message OR that the HIT Module incorporates/stores in the patient's laboratory result record a pointer to the exact data received in the LRI message.  Example: Placer Number; the HIT-originated Placer Number received in the LRI message may be incorporated/stored using a pointer rather than being stored redundantly in the patient's lab result record.	
S-EQ	Store equivalent	The HIT Module being tested must be designed to transform the exact data received in the LRI message to an equivalent format and then incorporate/store the equivalent format.  • Tester must verify that the HIT Module being tested transforms the exact data received in the LRI message to an equivalent format and incorporates/stores the equivalent format in the patient's laboratory result record.	
S-TR-R	Translate and store translation (exact value can be re-created from translation any time)	The HIT Module being tested must be designed to transform the exact data received in the LRI message to an equivalent value and then incorporate/store the equivalent value.  • Tester must verify that the HIT Module being tested incorporates/stores in the patient's laboratory result record the equivalent value.  • Tester must also verify that the HIT Module is able to re-create from this equivalent value the exact data received in the LRI message.	
S-RC	Process and re-create	The HIT Module being tested must be designed to process and incorporate/store in an "abstract-al manner" (e.g., using the HIT Module's data model) the exact data received in the LRI message and re-create the exact data (e.g., from the HIT Module's data model).  • Tester must verify that the HIT Module being tested processes and abstractly incorporates/stores in the patient's laboratory result record the exact data received in LRI message.  • Tester also must verify that the HIT Module is able to re-create the exact data received in the LRI message by abstracting the data (e.g., from the HIT Module's data model).  Example: Identifier Type Code; the HIT Module uses a separate file/table to store Social Security Numbers versus internal Medical Record Numbers, and does not need to retain the Identifier Type Code	