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## **Clinical Laboratory Inspection Checklist- Random**

Name of the Facility: _			
Date of Inspection:	/_	/_	

Ref.	Description	Yes	No	N/A	Remarks		
5	STANDARD ONE: REGISTRATION AND LICENSURE PROCEDURES						
5.5.	The health facility shall provide documented						
5.5.	evidence of the following:						
5.5.1.	Equipment maintenance services						
5.5.2.	Laundry services						
F F 2	Medical waste management as per Dubai						
5.5.3.	Municipality (DM) requirements						
5.5.4.	Housekeeping services						
	Calibration of temperature dependent equipment						
5.5.5.	(Refrigerator, freezer, incubator, water bath, room						
	temperature monitoring device etc.)						
5.5.6.	Calibration of centrifuges, weighing balance,						
3.3.0.	pipette, validation of biological safety cabinet.						
5.5.7.	Change of HEPA filter annually and fume hood						
5.5.7 .	validation.						
	The health facility shall maintain charter of						
5.7.	patients' rights and responsibilities posted at the						
5.7.	entrance of the premise in two languages (Arabic						
	and English).						
	The health facility shall ensure it has in place						
	adequate lighting and utilities, including						
5.8.	temperature controls, water taps, medical gases,						
	sinks and drains, lighting, electrical outlets and						
	communications.						

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5.11.	he Clinical Laboratory license shall be visibly
pc	osted at the reception of the health facility.
6 S1	TANDARD TWO: HEALTH FACILITY REQUIREMENTS
Al	Il DHA licenced Clinical Laboratories shall obtain
6.2. ac	ccreditation as per the DHA Clinical Laboratory
Ad	ccreditation Policy.
Th	he health facility should install and operate
eq	quipment required for provision of the proposed
6.4. se	ervices in accordance to the manufacturer's
sp	pecifications and should be validated for it's
in	tended use prior to using it in the service.
6.7.	he health facility design shall provide assurance of
0.7. pa	atients and staff safety.
7 S1	TANDARD THREE: HEALTHCARE PROFESSIONALS REQUIREMENTS
Al	II healthcare professionals in the DHA licensed
7.1.	ealth facility must hold an active DHA
	rofessional license and work within their scope of
pr	ractice.
Al	Il healthcare professionals directly dealing with
7.4.	atient should maintain a valid
	aining/certification of one (1) or more of the
fo	llowing:
7.4.1 Ba	asic Cardiopulmonary Resuscitation (CPR)
7.4.2 Ba	asic Life Support (BLS)
7.4.3 Ad	dvanced Cardiac Life Support (ACLS)
Th	ne Medical/Laboratory Director in an
ine	dependent clinical laboratory or laboratory in
ho	ospital setup shall be a full time DHA licensed
7.6.1. Pa	athologist (Clinical Pathologist or Anatomic
Pa	athologist) or a licensed holder of a doctoral
de	egree in chemistry, physical, biological, or clinical
lal	boratory science from accredited institution
763	case of a specialized laboratory, a licensed
7.6.2.	inical Laboratory Scientist (CLS) with doctoral

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	degree in the specialized field and appropriate				
	relevant training and experience may serve as the				
	Laboratory Director.				
	In an Outpatient Care Facility, a full time/part time				
7.6.3.	pathologist may be the designated laboratory				
	director.				
	To ensure safe and high-quality care is upheld				
	within DHA licensed facilities the Medical				
7.6.4.	Director/laboratory director shall abide by the				
	DHA Policy for Role and Responsibilities of Medical				
	Director.				
8	STANDARD FOUR: MANAGEMENT RESPONSIBIL	ITIES			
	Ensure all healthcare professionals employed have				
0.4.5	a current DHA license, are privileges as per the				
8.1.5.	Clinical Privileging Policy and work within their				
	scope of practice.				
	Maintain the recommended immunizations for				
	health professionals working at the clinical				
8.1.6.	laboratory, as per the DHA policy for Health				
	Screening and Immunization of Healthcare				
	Professionals.				
	Obtain prior approval from the Ministry of Health				
	and Prevention (MOHAP) for media and				
8.1.17.	advertisement materials, for further information				
0.1.17.	regarding the media and advertisement materials				
	approval procedures and requirements please visit				
	the MOHAP website.				
9	STANDARD FIVE: MANAGEMENT OF THE CLINIC	AL LABO	RATORY	OPERA	TIONS
9.1.1.	Requisition form				
	Should contain the patient's identity, age, sex,				
L	location, name of physician, last menstrual period,				
b.	date of specimen collection, source of specimen				
	when appropriate and the investigations requested.				
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9.1.2.	Receiving specimens from outside the country will require the licensed health facilities/clinical		
J	require the licensed health facilities/clinical		
	laboratories to obtain approval from MOHAP		
9.1.3.	Specimen collection		
	It can be done at the patient's bedside or in the		
	laboratory phlebotomy room/specimen collection		
b.	facilities depending on the type of specimen		
	required for the test.		
	•		
_	Physicians, nurses or medical laboratory		
e.	technologist can collect specimens who are		
	regularly trained, to ensure their competency.		
	Clinical Laboratory may have a "Primary Specimen		
	Collection Manual", containing information on		
	patient preparation before specimen collection (if		
	any), and exact methodology of specimen collection		
g.	(type of collection container and amount of		
	specimen to be collected, Phlebotomy order of		
	draw and instructions for fill volume and proper		
	mixing) labelling, handling, transportation and		
	storage of the specimens.		
	Specimens shall be secured appropriately to		
	prevent leakage, spillage or contamination. They		
I.	must be transported in biohazard bags and sent to		
h.	the laboratory along with the completed laboratory		
	requisition form (in the absence of electronic data		
	transfer).		
	A biohazard symbol shall be used on the specimen		
i.	transportation containers during transportation.		
	Appropriate specimen transportation kit (such as		
j.	use of dry ice, ice packs, etc.) shall be used		
	wherever required.		
	Clinical laboratories shall have clear criteria for		
k.	rejection of samples such as haemolysed or lipemic		
	samples.		

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9.1.4.	Accession List		
	Record of all the specimens received by the		
a.	laboratory for analysis shall be prepared by the		
	laboratory at the time of specimen receipt.		
	The accession list must record the patient's		
	identity including name, age, sex, location in the		
b.	hospital/health facility, name of referring physician,		
D.	investigations requested, date and time of receipt		
	of specimen and condition of the specimen at		
	receipt.		
	Assigns a unique number to register each specimen		
C.	received, which can be used to trace the specimen.		
C.	The test results and remarks if any shall also be		
	entered in the accession list.		
9.1.5.	Reporting test results		
	Test results approved and signed by the most		
a.	responsible physician/team and shall be made		
	available to the ordering physician.		
	Results shall be reported clearly, without any		
Ь.	errors, specifying measurement procedure where		
J.	appropriate and units of measurement. Test results		
	shall also contain the following:		
i.	Reference intervals as applicable		
ii.	Patient details with identification number		
iii.	Test details		
iv.	Date and time of specimen collection		
v.	Result reporting		
vi.	Specimen source		
vii.	Name		
IX.	Referral laboratory when applicable.		
	For quantitative test, laboratory shall not report		
C.	any numeric result outside the Analytical		
	Measurement Range of the analyser, unless the		 

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	sample is processed by dilution, a mixing procedure		
	or concentration.		
9.1.6.	Ethical considerations		
	Healthcare Professionals working in the clinical		
d.	laboratory shall not use expired		
	reagents/kits/specimen collection supplies		
	The clinical laboratory shall validate or verify assay		
i.	performance of new tests, methods, or instruments		
	prior to patient testing.		
ii.	Evidence of documented validation must be readily		
11.	available for any inspection.		
9.1.7.	Outsourcing Clinical Laboratory Services		
a.	The outsourced laboratory shall:		
	Be accredited as per the DHA Clinical Laboratory		
l.	Accreditation Policy.		
11	Have a primary sample collection and handling		
II.	manual, which may be electronic.		
	Clinical Laboratory test shall be performed only		
V.	upon a request from a DHA licensed Physician and		
٧.	sent along with the specimen to the testing Clinical		
	laboratory.		
	Clinical laboratories shall refrain from promoting or		
	marketing laboratory tests aiming to attract		
VI.	patients directly to visit the laboratory without		
	consulting or referring from the DHA licensed		
	treating Physician.		
9.1.8.	Outsourcing Clinical Laboratory Services outside		
3.1.0.	the UAE		
a.	It is prohibited to send patient's samples outside		
u.	UAE, unless the following conditions are met:		
l.	Unavailability of the requested medical test within		
14	the laboratory services in the UAE.		

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	Report from a consultant physician in case of rare		
II.	tumor whose specialization does not exist in the		
	UAE.		
III.	Sending samples should be through a laboratory or		
111,	health facility licensed by the DHA.		
	Clinical laboratories receiving the sample outside		
IV.	the country should be accredited; in accordance		
IV.	with the DHA Clinical Laboratory Accreditation		
	Policy.		
	Notify DHA with the entire laboratory tests sent		
V.	outside the UAE. Permission shall be obtained once		
	per specific laboratory test.		
9.2.1.	General Safety Considerations		
	Eyewash facility shall be available as "stand-alone"		
	facility or attached to sink or portable. Sealed		
	single use solution bottles may also be used. At		
	locations where, hazardous chemicals are handled		
	by employees, emergency eyewash and shower		
III.	equipment shall be available no greater than ten		
	(10) seconds travel distance from areas in the		
	laboratory where hazardous chemicals are present		
	with unobstructed pathway. The door must be		
	open in the direction toward the eyewash/ shower		
	station.		
15.7	Laboratories shall ensure proper preservation and		
IV.	security of specimens.		
1/1	Periodic checking of all safety equipment and		
VI.	accessories shall be ensured.		
	For reasons of both safety and security, personal		
	belongings (coats, bags, pocketbooks, etc.) must		
IX.	not be kept in the work areas of the laboratories.		
	Personal belongings must be secured in employees'		
	lockers or staff designated areas.		
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V	The laboratory shall restrict the access to testing		
X.	area; for authorized personnel only.		
9.2.2.	Biohazard Materials		
	An updated list of hazardous materials used in the		
	laboratory shall be maintained. All hazardous		
a.	materials shall be accounted for on a continuous		
	basis.		
	Biohazard symbol shall be used on all containers		
b.	containing biohazard materials while being		
	transported to the laboratory or disposed of.		
f.	All anatomic pathology wastes are placed in a		
1.	biohazard waste container for incineration.		
ď	Biohazard spill kits and chemical spill kits must be		
g.	available in the Laboratory.		
9.2.3.	Chemical Safety		
b.	Chemical Safety Data Sheet (SDS) shall be		
D.	available and accessible to all staff.		
C.	Containers of hazardous chemicals shall have		
C.	precautionary labels indicating type of hazard.		
	The laboratory shall limit the storage of flammable		
d.	and combustible chemicals as per the amount		
u.	required and shall store these chemicals inside		
	flammable storage cabinet.		
9.2.4.	Handling Sharps		
	Recapping of needles is strictly prohibited.		
	Contaminated needles or other sharps must not be		
	sheared, bent, recapped, or removed from syringes		
	or other devices unless it can be accomplished		
a.	using a mechanical device (such as a haemostat) or		
	by using re-sheathing instruments or self-		
	sheathing needles or retractable needles with		
	locking system to prevent recapping of needles by		
	hand. The laboratory shall have procedures to		

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	follow after a sharp injury; that includes needlestick			
	injuries.			
	Sharps (i.e., needles, syringes with attached			
	needles, scalpel blades) must be placed in a stable,			
	rigid, puncture-resistant "sharps" container labelled			
Ь.	with a biohazard warning label. Slides, coverslips,			
	and capillary tubes may be placed in a rigid,			
	puncture-resistant container or red-bagged			
	biohazard waste container.			
	Sharps containers must not be overfilled. When a			
C.	sharps container becomes two-thirds full, seal and			
C.	discard it into a red-bagged biohazard waste			
	container or into a red bag for incineration.			
9.2.5.	Fire Safety			
	Train staff to respond to fire events on the			
g.	premises.			
9.3.	Health Records			
	Laboratories sending reports electronically should			
	include electronic signature of the authorized			
9.3.3.	signatory. Laboratories should be able to provide			
	critical information required by a physician on			
	telephone.			
9.3.5.	Equipment maintenance reports must be kept for			
9.5.5.	future reference.			
	Patient Result Records and materials shall be			
9.3.6.	retained aligned to the DHA Guidelines for			
	Managing Health Records.			
10	STANDARD SIX: MOBILE LABORATORIES			
10.2.4	The mobile laboratory should be accredited as per			
10.2.1.	the Clinical Laboratory Accreditation Policy.			
	The scope of services provided by the mobile			
4005	laboratory should be elaborated and documented			
10.2.5.	and the mobile laboratory should not provide any			
		•	i	i e

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	laboratory licensed to provide phlebotomy services,			
	shall only provide that service).			
	There should be a dedicated space to park the			
10.2.6.	vehicle to ensure the temperature requirement in			
	the vehicle is maintained, as required.			
	Approved vendors shall maintain all the equipment			
10.2.8.	and the maintenance performed shall be			
	documented.			
	Equipment that is very sensitive to movement and			
10.2.9.	fluctuation in temperature shall not be used in a			
	mobile laboratory.			
	All healthcare professionals should be employed as			
10.2.11.	per the service provided and should maintain a			
10.2.11.	valid training/certification in basic CPR, or BLS or			
	ACLS.			
11	STANDARD SEVEN: WALK-IN LABORATORY TES	TS		
	Licensed Clinical Laboratories may accept walk-in			
11 1	patients for specific laboratory tests that do not			
11.1.				
11.1.	patients for specific laboratory tests that do not			
	patients for specific laboratory tests that do not require a physician's order. Refer to the list of			
11.1. 11.1.1.	patients for specific laboratory tests that do not require a physician's order. Refer to the list of services permitted elaborated in <b>Appendix 3</b> .			
	patients for specific laboratory tests that do not require a physician's order. Refer to the list of services permitted elaborated in <b>Appendix 3</b> .  Critical laboratory results shall be reported clearly	SION SEI	RVICES	
11.1.1.	patients for specific laboratory tests that do not require a physician's order. Refer to the list of services permitted elaborated in <b>Appendix 3</b> .  Critical laboratory results shall be reported clearly to the patient.	SION SEI	RVICES	
11.1.1.	patients for specific laboratory tests that do not require a physician's order. Refer to the list of services permitted elaborated in <b>Appendix 3</b> .  Critical laboratory results shall be reported clearly to the patient.  STANDARD EIGHT: BLOOD BANK AND TRANSFU	SION SEI	RVICES	
11.1.1. 12	patients for specific laboratory tests that do not require a physician's order. Refer to the list of services permitted elaborated in <b>Appendix 3</b> .  Critical laboratory results shall be reported clearly to the patient.  STANDARD EIGHT: BLOOD BANK AND TRANSFU Blood bank and Transfusion Service shall have a	SION SEI	RVICES	
11.1.1. 12	patients for specific laboratory tests that do not require a physician's order. Refer to the list of services permitted elaborated in <b>Appendix 3</b> .  Critical laboratory results shall be reported clearly to the patient.  STANDARD EIGHT: BLOOD BANK AND TRANSFU Blood bank and Transfusion Service shall have a medical director who is a licensed physician,	SION SEI	RVICES	
11.1.1. 12	patients for specific laboratory tests that do not require a physician's order. Refer to the list of services permitted elaborated in <b>Appendix 3</b> .  Critical laboratory results shall be reported clearly to the patient. <b>STANDARD EIGHT: BLOOD BANK AND TRANSFU</b> Blood bank and Transfusion Service shall have a medical director who is a licensed physician, qualified by training, experience.	SION SEI	RVICES	
11.1.1. 12	patients for specific laboratory tests that do not require a physician's order. Refer to the list of services permitted elaborated in <b>Appendix 3</b> .  Critical laboratory results shall be reported clearly to the patient. <b>STANDARD EIGHT: BLOOD BANK AND TRANSFU</b> Blood bank and Transfusion Service shall have a medical director who is a licensed physician, qualified by training, experience.  Blood & components storage devices shall have the	SION SEI	RVICES	
11.1.1.  12  12.1.	patients for specific laboratory tests that do not require a physician's order. Refer to the list of services permitted elaborated in <b>Appendix 3</b> .  Critical laboratory results shall be reported clearly to the patient.  STANDARD EIGHT: BLOOD BANK AND TRANSFUR Blood bank and Transfusion Service shall have a medical director who is a licensed physician, qualified by training, experience.  Blood & components storage devices shall have the capacity and design to ensure that the proper	SION SEI	RVICES	
11.1.1.  12  12.1.	patients for specific laboratory tests that do not require a physician's order. Refer to the list of services permitted elaborated in <b>Appendix 3</b> .  Critical laboratory results shall be reported clearly to the patient. <b>STANDARD EIGHT: BLOOD BANK AND TRANSFU</b> Blood bank and Transfusion Service shall have a medical director who is a licensed physician, qualified by training, experience.  Blood & components storage devices shall have the capacity and design to ensure that the proper temperature is maintained, monitored for storage	SION SEI	RVICES	
11.1.1.  12  12.1.	patients for specific laboratory tests that do not require a physician's order. Refer to the list of services permitted elaborated in <b>Appendix 3</b> .  Critical laboratory results shall be reported clearly to the patient. <b>STANDARD EIGHT: BLOOD BANK AND TRANSFU</b> Blood bank and Transfusion Service shall have a medical director who is a licensed physician, qualified by training, experience.  Blood & components storage devices shall have the capacity and design to ensure that the proper temperature is maintained, monitored for storage of blood & component.	SION SEI	RVICES	

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	blood, blood components, reach unacceptable			
	conditions			
13	STANDARD NINE: CYTOGENETIC TESTING SERVI	CES		
	Genetic testing shall only be requested by a DHA			
13.1.	licensed Physician with enough clinical justification,			
15.1.	after patient consultation, in a DHA licensed Health			
	facility.			
13.3.	A Consultant/Specialist Physician shall be			
13.3.	consulted before and after genetic testing.			
	Only the referring/treating Physician shall make			
13.5.	recommendations or prescribe to the patient any			
13.3.	medication or healthcare products, based on the			
	laboratory results.			
	The Cytogenetic laboratory records and results			
13.9.	shall accurately reflect all stages of the process and			
	all results obtained.			
13.10.	The Cytogenetic laboratory records shall include			
15.10.	the following:			
13.10.1.	Media used			
13.10.2.	Reactions observed			
13.10.3.	Number of cells counted			
13.10.4.	Number of cells karyotyped			
13.10.5.	Number of chromosomes counted for each			
13.10.3.	metaphase spread			
13.10.6.	Quality of the banding.			
14	STANDARD TEN: MOLECULAR TESTING SERVICE	S		
	Validation studies shall include representatives			
14.2.	from each specimen type expected those that are			
14.2.	to be tested in the assay and specimens			
	representing the scope of reportable results.			
	Molecular testing reports shall include specific		_	
14.3.	testing information including the following			
	information:			

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14.3.1.	Testing methodology			
14.3.2.	Limitations of the method			
14.3.3.	Interpretation of findings			
14.3.4.	Recommendations for additional testing.			
15	STANDARD ELEVEN: MOLECULAR GENETIC SERV	ICES		
15.1.	Molecular Genetic testing shall only be carried out			
15.1.	against a DHA licensed Physician's order.			
	The DHA licensed health facility shall refrain from			
15.2.	requesting or performing paternity or genealogical			
	tests; except in the following cases:			
15.2.1.	Obtain prior official written approval from DHA.			
	Upon a request from the competent judicial			
15.2.2.	authorities in UAE and the concerned health			
	authorities.			
15.3.	All tests carried out should be FDA approved or			
	equivalent.			
15.6.	Molecular genetic testing reports shall include the			
45.64	following information:			
15.6.1.	List of mutant genes for alleles tested.			
15.6.2.	Any recommendations for referral to a genetic counsellor.			
15.6.3.	Detection rate of the test.			
15.0.5.				
15.6.4.	Use of standard nomenclature for genes and mutations.			
15.6.5.	Clinical implications of mutations detected.			
15.0.5.	The laboratory should consider three categories of			
15.7.	test performance in the evaluation process:			
15.7.1.	Analytic validity			
15.7.2.	Clinical validity			
15.7.3.	Clinical utility.			
16	STANDARD TWELVE: CYTOPATHOLOGY SERVICE	-s		
	Each individual evaluating cytology preparations by			
16.2.	manual microscopic technique shall examine no			
	mandai microscopic technique shall examille no			

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	more than hundred (100) slides (gynaecologic and			
	non-gynaecological or both) in a day.			
	For the laboratory that perform immunochemical			
	tests that provide predictive information that are			
	independent of diagnosis or other cytopathologic			
16.3.	findings, the patient report must include			
	information on specimen fixation, specimen			
	processing, antibody clone used and the scoring			
	method used.			
16.7.	All cytopathologic reports shall be authorized by a			
10.7.	DHA licensed pathologist.			
	The cytopathology reports shall include a concise			
16.8.	descriptive diagnosis in a standard descriptive			
10.0.	terminology that includes a general categorization			
	and descriptive diagnosis.			
	The laboratory shall promptly notify the			
16.9.	responsible clinician(s) when there are changes in			
	the reports that significantly affect patient care.			
	Cytology slides and blocks are properly stored in a			
	temperature controlled, pest-free, organized			
16.10.	manner to prevent contamination from blood or			
	other fluids or tissue and be readily accessible for			
	retrieval.			
17	STANDARD THIRTEEN: HISTOPATHOLOGY SERV	/ICES	1	
	All macroscopic tissue gross examinations are			
	performed by a DHA licensed pathologist or by			
17.1.	qualified competent laboratory DHA licensed			
	healthcare professional under the supervision of a			
	qualified pathologist.			
17.3.	All histopathology results shall be reviewed and			
	authorized by a DHA licenced pathologist.			
17.5.	The laboratory shall promptly notify the			
	responsible clinician(s) when there are changes to			

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	reports such as that significantly affect patient				
	care.				
17.6	The laboratory shall correlate the results of				
17.6.	specialized studies with the morphologic diagnosis.				
	Slides and blocks shall be stored properly in a				
	temperature-controlled, pest-free, organized				
17.7.	manner to prevent contamination from blood or				
	other fluids or tissues and be readily accessible for				
	retrieval.				
	When frozen section and final diagnosis results are				
17.10.	discrepant, there is a review of findings, and the				
17.10.	discrepancy is resolved and shall record this in the				
	final report.				
18	STANDARD FOURTEEN: ALCOHOL AND DRUG TE	ESTING S	ERVICES	;	
	Clinical Laboratories should apply for alcohol, drug				
18.1.	and substance abuse testing through Sheryan				
	(under amend facility license request).				
	Alcohol and Drug Testing Laboratories should				
18.2.	acquire the following accreditations within 2 years				
	of application:				
18.2.1.	College of American Pathologists (CAP)				
18.2.2.	ISO accreditations ISO/IEC 17025				
	Each laboratory should have the capability to				
18.4.	perform the screening and confirmatory tests at				
	the same laboratory site.				
	If confirmatory test is not done in the same clinical				
	laboratory, then there should be a contract in place				
18.4.1.	with the outsourced laboratory that clearly includes				
	the name of laboratory and their accreditation on				
	each tested drug/substance.				
	The testing procedure of each laboratory shall be				
18.5.	capable of detecting drugs, drugs metabolites,				
	adulterants, and substituted specimens.				

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	The specimen should be treated as evidence and all				
18.6.	aspects of the procedure shall be documented and				
	available for possible court testimony.				
	The availability of MOHAP approved Evidential				
18.8.	Breath analyser is mandatory to conduct alcohol				
	testing.				
	The analyser should have a distinguishing level for				
18.8.1.	alcohol from acetone at the 0.02% alcohol				
	concentration.				
	The laboratory shall have the capability of				
	conducting blood alcohol concentration, or urine				
18.9.	alcohol level, these tests can be used as an extra				
	evidential method in cases of positive breath				
	analyser test.				
	An additional approval from General Civil Aviation				
18.10.	Authority is required to conduct Alcohol and Drug				
	Testing for all aviation industries.				
					l.
19	STANDARD FIFTEEN: DRIVE THROUGH LABORAT	TORY SEI	RVICES	1	
19	STANDARD FIFTEEN: DRIVE THROUGH LABORATE The drive through laboratory services shall be	TORY SEI	RVICES		
<b>19</b> 19.2.		TORY SEI	RVICES		
	The drive through laboratory services shall be	TORY SEI	RVICES		
	The drive through laboratory services shall be provided in an area within the premise of a DHA	TORY SEI	RVICES		
	The drive through laboratory services shall be provided in an area within the premise of a DHA licensed Health facility.	TORY SEI	RVICES		
19.2.	The drive through laboratory services shall be provided in an area within the premise of a DHA licensed Health facility.  The drive through laboratory services shall be	FORY SEI	RVICES		
19.2.	The drive through laboratory services shall be provided in an area within the premise of a DHA licensed Health facility.  The drive through laboratory services shall be provided only by DHA licensed healthcare	TORY SEI	RVICES		
19.2. 19.5.	The drive through laboratory services shall be provided in an area within the premise of a DHA licensed Health facility.  The drive through laboratory services shall be provided only by DHA licensed healthcare professionals (Registered Nurses or Phlebotomists)	TORY SEI	RVICES		
19.2.	The drive through laboratory services shall be provided in an area within the premise of a DHA licensed Health facility.  The drive through laboratory services shall be provided only by DHA licensed healthcare professionals (Registered Nurses or Phlebotomists)  Testing shall be initiated only upon physician's	TORY SEI	RVICES		
19.2. 19.5.	The drive through laboratory services shall be provided in an area within the premise of a DHA licensed Health facility.  The drive through laboratory services shall be provided only by DHA licensed healthcare professionals (Registered Nurses or Phlebotomists)  Testing shall be initiated only upon physician's order and the result should be sent back to the	TORY SEI	RVICES		
19.2. 19.5. 19.6.	The drive through laboratory services shall be provided in an area within the premise of a DHA licensed Health facility.  The drive through laboratory services shall be provided only by DHA licensed healthcare professionals (Registered Nurses or Phlebotomists)  Testing shall be initiated only upon physician's order and the result should be sent back to the treating physician. A copy of the results could be	FORY SEI	RVICES		
19.2. 19.5.	The drive through laboratory services shall be provided in an area within the premise of a DHA licensed Health facility.  The drive through laboratory services shall be provided only by DHA licensed healthcare professionals (Registered Nurses or Phlebotomists)  Testing shall be initiated only upon physician's order and the result should be sent back to the treating physician. A copy of the results could be shared with the patient.	TORY SEI	RVICES		
19.2. 19.5. 19.6.	The drive through laboratory services shall be provided in an area within the premise of a DHA licensed Health facility.  The drive through laboratory services shall be provided only by DHA licensed healthcare professionals (Registered Nurses or Phlebotomists)  Testing shall be initiated only upon physician's order and the result should be sent back to the treating physician. A copy of the results could be shared with the patient.  All staff shall maintain a current Basic Life Support	TORY SEI	RVICES		
19.2. 19.5. 19.6.	The drive through laboratory services shall be provided in an area within the premise of a DHA licensed Health facility.  The drive through laboratory services shall be provided only by DHA licensed healthcare professionals (Registered Nurses or Phlebotomists)  Testing shall be initiated only upon physician's order and the result should be sent back to the treating physician. A copy of the results could be shared with the patient.  All staff shall maintain a current Basic Life Support (BLS) certification.	TORY SEI	RVICES		
19.2. 19.5. 19.6.	The drive through laboratory services shall be provided in an area within the premise of a DHA licensed Health facility.  The drive through laboratory services shall be provided only by DHA licensed healthcare professionals (Registered Nurses or Phlebotomists)  Testing shall be initiated only upon physician's order and the result should be sent back to the treating physician. A copy of the results could be shared with the patient.  All staff shall maintain a current Basic Life Support (BLS) certification.  The drive through laboratory facility shall maintain	TORY SEI	RVICES		

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	immediate and safe provision of care, if required.				
	Mock drill should be conducted accordingly				
	Tests that requires special phlebotomy procedure				
19.10.	such as blood culture tests and which requires a				
19.10.	sterile technique, shall not be performed at these				
	facilities.				
	These facilities shall have a tunnel for entrance of				
	the vehicle, where appropriate temperature is				
19.11.	maintained, a registration area, and a dedicated				
	storage area with maximum temperatures of 24				
	degrees centigrade.				
	There shall be availability of an adjustable table to				
19.13.	be inserted through the door of the car, in order for				
	the patient to rest the arm.				
	Ensure provision of a blood collection chair for				
19.14.	patients from who blood cannot be collected while				
	sitting in the car.				
19.17.	Patients privacy should be ensured, especially for				
19.17.	female patients.				
	Infants and children below the age of seven (7)				
19.18.	years shall not tested at a drive through laboratory				
	services.				
20	STANDARD SIXTEEN: SAMPLE COLLECTION SITE				
20.4	Sample collection site shall require a separate DHA				
20.1.	license as a Medical laboratory.				
	The collection site shall be divided into minimum				
20.4.	sections of waiting, collection, receiving, storage				
	and medical waste.				
20.5.	Patient privacy shall be maintained at all times.				
	There shall be a minimum of two full-time				
20.6	healthcare professionals in the collection site who				
20.6.	may be a DHA licensed physician, registered nurse				
	or phlebotomists.				
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20.7.	The collection site shall obtain accreditation within two (2) years from the date of obtaining the license and upon renewal.		
APPENDIX 2:	GUIDELINES ON QUALITY ASSURANCE		
4	Practice of IQC includes the following:		
	For quantitative tests, laboratories should perform		
4.4	IQC every day by using 2 levels of QC (high and		
4.4.	low) on tests run daily or every time the tests are		
	run in case of infrequently run tests.		
5	The level of QC per day for Quantitative Tests:		
	The following guidelines will be useful to the		
5.1.	laboratories in the practice of IQC using either one		
	level or two-level QC materials:		
5.1.1.	When one level QC is used: Reject test run if		
3.1.1.	following errors occur:		
a.	Value is outside 3 SD (13s)		
b.	2 consecutive values are outside 2 SD on the same		
D.	side, but within 3 SD (22s)		
C.	4 consecutive value are outside 1SD on the same		
<b>C.</b>	side, but within 2SD (41s)		
5.1.2.	When two level QC are used: Reject test run if		
J.1.2.	following errors occur:		
a.	Either QC value is outside 3 SD (13s)		
b.	Both QC values are outside 2 SD on the same side,		
D.	but within 3SD (22s)		
	Difference between the two-level QC values is >4		
C.	SD i.e. one level QC is >2 SD and other level QC is <		
	2 SD (R4s)		
d.	10 consecutive values of the same level QC are		
	above or below the mean, but within 2 SD (10x)		
	5 consecutive values of one level QC and 5		
e.	consecutive values of the other level QC are above		
	or below the mean, but within 2 SD (10x).		

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APPENDIX 3:	PERMITTED WALK IN LABORATORY TESTS		
A3.1.	Anatomic Pathology: All tests under this category		
	requires Prescription		
	Clinical Biochemistry: General Clinical Biochemistry		
	Testing Does not require Prescription // Special		
A3.2.	Clinical Biochemistry testing such as body fluids		
	samples other than blood, especial enzymes and iso		
	enzymes require Prescription		
42.2	Therapeutic Drug Monitoring: All tests under this		
A3.3.	category requires Prescription		
A3.4.	Drug Abuse Monitoring: All tests under this		
	category requires Prescription		
	Haematology: General haematology testing Does		
A3.5.	not require Prescription// Special haematology		
	testing such as Malaria film, Bone marrow smear		
	and biopsy require Prescription		
	Coagulation: General Coagulation testing Does not		
A3.6.	require Prescription// Special Coagulation testing		
A3.6.	such as Factor testing, protein C, Protein S and		
	Lupus Anticoagulant require Prescription		
	Microbiology: General Microbiology and culture Do		
	not require Prescription// Special Microbiology		
A3.7.	testing such as blood cultures, tissue sample		
	cultures, CSF cultures and cultures which requires		
	special sample to be collected require Prescription		
A3.8.	Immunology and endocrinology: General		
	immunology and endocrinology Do not require		
	Prescription// Special immunology and		
	endocrinology testing such as hormonal		
	suppression and challenging tests require		
	Prescription		
A3.9.	Infectious disease: Permitted walk in laboratory		
AJ.3.	tests: Infectious disease screening on condition		

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	that laboratory must comply with patients consent		
	and notification of positive cases for authorities		
A3.10.	Blood bank and Transfusion: Permitted walk in		
	laboratory tests: Only blood grouping for screening		
	purposes and not for blood transfusion// All other		
	tests under this category requires Prescription		
A3.11.	Histocompatibility (HLA lab): All tests under this		
	category requires Prescription		
A3.12.	Molecular biology: All screening molecular biology		
	testing such as Respiratory, GIT, STDs panels, and		
	HPV screening Do not require Prescription// All		
	other tests under this category requires		
	prescription		
A3.13.	Genetics: All tests under this category requires		
	Prescription		
A3.14.	Forensic pathology: All tests under this category		
	requires prescription		

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