

# OSBI Drug Laboratory Training Manual Revision #19, Effective Date 10-20-2025

### **Table of Contents**

Introduction	3
Orientation to the OSBI	2
General Knowledge of Forensic Science	4
Applicable Criminal and Civil Law and Procedures	4
Drug Quality System Overview	5
Miscellaneous	5
Testimony & Presentation of Evidence in Court	6
Weights and Measures Utilized in Drug and Marijuana Reports and Balance Scale Calibration and Uncertainty (DR-3 and DR-4)	7
Cannabis Analysis	12
Drug Literature	16
Pharmaceutical Identification by Literary Reference (DR-5)	18
Training Procedures for Extractions & Handling	20
Identification of Gamma-Hydroxybutyric Acid through Derivatization (DR-7)	23
Training Procedures for Color Tests (DR-10, DR-11, DR-13)	25
Cocaine Free-base Determination by Hexane Solubility (DR-21)	27
Identification of Lysergic Acid Diethylamide (DR-101)	29
Gas Chromatography Analysis (Flame Ionizing Detector) (DR-30)	30
Analysis of Mushrooms to Determine the Presence of Psilocyn or Psilocybin (DR-45)	33
PDF Examination Documentation Procedure (DR-50)	35
Drug Analysis by FTIR (DR-60)	37
Requirements Prior to Drug Analysis Using FTIR	41
Gas Chromatograph Mass Spectrometer Methods for Drug Analysis (DR-70)	42
Administrative and Technical Reviewing of Casework	46
Requirements Prior to Drug Analysis	47
Liquid Nitrogen Safety	49
Gas Chromatograph Infrared Detector Methods for Drug Analysis (DR-75)	51
Requirements Prior to Drug Analysis using the GCIRD	52



# OSBI Drug Laboratory Training Manual Revision #19, Effective Date 10-20-2025

Controlled Substances Technician	53
Appendix I - Basic Operation of the Gas Chromatograph	56
Appendix II - Basic Operation of the Mass Spectrometer	58
Appendix III – Helpful Tips for The Mass Spectrometer	63
Appendix IV - Mass Peaks of Common Contaminants	66
Appendix V – Isomers	67
Mock Trial Evaluation Form	69
Additional Reading/Training	70
Approval	71
History	72



Revision #19, Effective Date 10-20-2025

#### Introduction

The Controlled Substances Laboratory of the Criminalistics Services Division (CSD) of the Oklahoma State Bureau of Investigation (OSBI) is part of an accredited full-service laboratory system responsible for the analysis of samples suspected to contain a controlled dangerous substance. This training manual is intended to provide an analyst with the skills and information needed to perform analysis of submitted samples. Each section of this manual lists a specific goal and the tasks that a trainee should complete in order to achieve this goal. The training will be assessed using an oral examination as well as a competency examination.

At the conclusion of training the trainee should have the following:

- 1. Knowledge of the principles and practices of forensic marijuana and drug analysis as they relate to the analysis of case material.
- 2. Knowledge of the theory and application of instrumentation and specialized techniques used to examine marijuana, controlled substances, and non-controlled substances.
- The skills and ability to perform accurate forensic analysis independently and proficiently, to accurately document the findings of all analysis in accordance with the appropriate policies and procedures, and to accurately generate a report on those findings.

If an analyst has previously passed a mock trial in the OSBI Controlled Substances Unit and has previous experience testifying in court, the analyst may be given an oral examination in lieu of a second mock trial. Two or more Senior Criminalists and the Technical Manager or designee will be present during the mock trial and the "Mock Trial Evaluation Form" will be used for grading. Requirements for passing include a minimum score of a 2 for each section and approval from the Technical Manager. Any score lower than a 2 must have a written explanation for the score.

This training manual can be modified by the Technical Manager for re-training purposes, including an analyst that is returning to the Controlled Substances Unit from another discipline or an analyst that needs retraining in a specific area for remedial reasons.

Once released for casework, it is up to the analyst to seek further training for the maintenance of skills and expertise. The Technical Manager will periodically send out articles for the analyst to read; those articles are to be documented on the Additional Reading/Training form.

The finalized training notebook will be kept by the analyst at the OSBI laboratory. Once training is completed, the training notebook will be scanned and uploaded into the analyst's individual folder on the QA server. Upon termination or transfer to another unit, the training notebook will be scanned, if not already in digital format, and uploaded into the analyst's individual folder on the QA server.



Revision #19, Effective Date 10-20-2025

#### **Orientation to the OSBI**

#### Goals

- To ensure the trainee is familiar in forensic science and the different types of forensic services
- To familiarize the trainee with criminal and civil laws that pertain to forensic chemistry
- To familiarize the trainee with the OSBI Drug Quality System
- To introduce the trainee to the OSBI laboratory management systems
- To introduce the trainee to courtroom testimony dynamics and behavior in the courtroom

### **General Knowledge of Forensic Science**

#### Literature

Date	Literature
	Smith, F.P. Overview of Forensic Drug Analysis. Handbook of Forensic Drug
	Analysis. 2005, pages 1-12

### **Applicable Criminal and Civil Law and Procedures**

#### Literature

Date	Literature
	Oklahoma State Statute Title 63, Chapter 2 – Uniform Controlled Dangerous
	Substances Act, www.oklegislature.gov
	Federal Code of Regulations Title 21 Part 1308
	Haggerty II, M.D. Confrontation and the Preliminary Hearing. Q & A: The
	Newsletter of the Criminal Law Section. Vol. 4, Issue 3, May-June 2006, pages 23-31
	Woodson, M. Relevance and Reliability: What All Expert Testimony Needs.
	Oklahoma Bar Journal, 79 OBJ 534, March 2008
	Calhoun, M. C. Scientific Evidence in Court: Daubert or Frye, 15 Years Later. Legal
	Backgrounder, Vol. 23, No. 37, August 22, 2008

Date	Tasks
	Discuss differences in distribution vs. trafficking vs. possession charges
	Discuss why drugs are scheduled and criteria for different schedules
	Discuss how drug laws are enacted, by vote of the people & legislative process,
	including the process of how drugs are controlled.



# OSBI Drug Laboratory Training Manual Revision #19, Effective Date 10-20-2025

### **Drug Quality System Overview**

#### Literature

Date	Literature/Tasks
	OSBI Controlled Substances Quality Assurance Manual
	Review OSBI CSD QM 7.4
	Review OSBI CSD QM 13, 14.1, 14.2, and 14.3
	Read Physical Evidence Quality Procedures Manual 2.1 Evidence Handling

### Discussion

#### Miscellaneous

#### Literature

Date	Literature/Tasks
	Review OSBI CSD QMA 2, Evidence Management Requirements, and observe
	evidence submitting procedures
	Review OSBI CSD QMA 3, Evidence Packaging and Sealing Guidelines, and observe
	evidence sealing and handling procedures

#### **Tasks**

The analyst will be shown where to locate the BEAST LIMS system and a brief
overview will be given
The analyst will observe checking out, inventorying and analyzing of a minimum of
5 cases
The analyst will be shown where to locate Chemical Inventory and a brief overview
will be given



Revision #19, Effective Date 10-20-2025

### **Testimony & Presentation of Evidence in Court**

#### Literature

Date	Literature
	Review OSBI Policy 108
	Shelton Hon., D.E., Barak, G., Kim, Y.S. A Study of Juror Expectation and Demands
	Concerning Scientific Evidence: Does the "CSI Effect" Exist. Selected Works
	(www.works.bepress.com). February 2007, pages 331-368

#### Discussion

Date	Tasks
	Discuss courtroom testimony and presentation of evidence with trainer
	Discuss bringing and opening evidence in court
	Discuss requirements for external testing requested by defense, including any accreditation requirements

#### Tasks

Review documentation when leaving evidence in court and how to enter the information into the BEAST
Review a Testimony Review form and Qualified Reviewer form
Review testimony report and who to send it to
Observe a Criminalist from the Controlled Substances Unit giving testimony

Approval	
Trainee	Date
Trainer/ Supervisor	Date
Comments	



Revision #19, Effective Date 10-20-2025

# Weights and Measures Utilized in Drug and Marijuana Reports and Balance Scale Calibration and Uncertainty (DR-3 and DR-4)

#### Goals

- To establish uniform guidelines in the determination and reporting of weights and volumes of substances submitted for analysis.
- To establish guidelines for procedures to document balance verification, calibration and uncertainty.
- To provide an understanding of the theory of uncertainty of measurement.
- To understand how uncertainty of measurement is calculated and factors that can affect uncertainty.
- To be able to explain uncertainty of measurement in a way a layperson can understand.

#### Literature

Date	Literature	
	PowerPoint Presentation for Uncertainty of Measurement	
	Bell, S. A Beginner's Guide to Uncertainty of Measurement. Measurement Good	
	Practice Guide No. 11 (Issue 2). National Physical Laboratory (PDF: Uncertainty of	
	Measurement)	
	Protocol DR-3 (Weights and Measures Utilized in Drug and Marijuana Reports)	
	Protocol DR-4 (Balance/Scale Calibration and Uncertainty)	
	Protocol DR-4 Attachment 1 (Budget for Calculating Uncertainty of Measurement)	
	Protocol DR-4 Attachment 2 (Controlled Substances Scale Scenarios)	
	Weighing the Right Way. Guide Book Proper Weighing with Laboratory Balances.	
	Mettler Toledo, 05/2012 (PDF: Uncertainty of Measurement)	
	M3003 The Expression of Uncertainty and Confidence in Measurement. United	
	Kingdom Accreditation Service, Edition 2, January 2007	

Date	Tasks	
	The definitions for: net weight, gross weight, approximate volume and residue	
	When gross weights can be utilized	
	The recommended report wording concerning significant digits for the reported	
	weight ranges	
	When a balance is to be checked for proper calibration and when a balance is to be	
	calibrated	
	The acceptable operating range of a balance	



Revision #19, Effective Date 10-20-2025

The proper procedure if a balance is not operating within specified operating range,
or if a balance is out of service
The recommended procedure for checking balances while working cases involving
trafficking charges
The reason for not reporting weights under a predetermined weight for the
different scales
Uncertainty of Measurement
The definition of Uncertainty of Measurement
How uncertainty of measurement is calculated and the factors considered,
including budget items vs items not included in budget
How often it is calculated and why it is recalculated
Proper report wording of uncertainty and when it is reported
Where should uncertainty be added to the report for marijuana cases that include
both grams and pounds
Trafficking levels
Trafficking weights for controlled substances (i.e., marijuana, meth, cocaine,
cocaine base, etc.)

#### **Tasks**

**Approval** 

Date	Tasks	
	Verify a bench top balance and record on appropriate OSBI DR4 form	
	Verify a large capacity scale and record on appropriate OSBI DR4 form	
	Verify an analytical balance and record on appropriate OSBI DR4 form	
	Perform 31-day Measurement Assurance Program for bench top balance	
	Perform 31-day Measurement Assurance Program for large capacity scale	
	Perform 31-day Measurement Assurance Program for analytical balance	
	Demonstrate how to determine the approximate volume of a container using the formula $V=\pi r^2h$	

Trainee	Date	
Trainer/ Supervisor	Data	
Supervisor	Date	
Comments		



Revision #19, Effective Date 10-20-2025

#### **Thermometers**

#### Goals

• To establish a guide for the proper reading of thermometers used in the laboratory

#### How to Read a Thermometer

Thermometers should be handled carefully because they are tubes of glass filled with either mercury or colored spirits.

Laboratory thermometers should NOT be shaken like the home variety thermometer. To lower the temperature, just cool them in a refrigerator or water/ice bath. Usually, they are either partial or whole immersion thermometers; this means that the bulb may be either partially submerged in a liquid or must be totally submerged in a liquid to accurately register the temperature. Thermometers used in the refrigerators are not to be submerged or placed into any liquid.

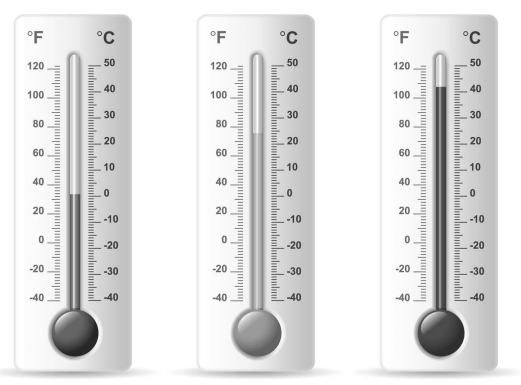
Place the thermometer in the material/refrigerator in which the temperature is to be measured. If you are measuring the temperature of a material while it is being heated, make certain that you do not let the thermometer rest on the bottom of the container and that the bulb is submerged in the material itself.

To read the temperature indicated on a thermometer, your eye should be at the level of the liquid in the thermometer. Read the thermometer to the appropriate number of digits. For example, a thermometer on which the heavy or extended lines are marked 10, 20, 30... should be read to the nearest degree.



Revision #19, Effective Date 10-20-2025

First examine the scales below, each degree is divided into smaller divisions. The number of divisions may vary between thermometers, so it is important to look at the scale.



Record the temperature of the three thermometers, to the nearest degree Celsius.

#1	#2	#3	

#### Literature

Date	Literature
	Read Physical Evidence Quality Procedures Manual 2.4, Evidence Refrigerator and
	Freezer Maintenance

Date	Tasks	
	What is to be done in the event the refrigerator/freezer is out of temperature	
	range?	
	When should monitoring the temperature be performed and how is this	
	documented?	



Revision #19, Effective Date 10-20-2025

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Date	Tasks	
Backup System or Generator		
	Trainer will demonstrate the proper steps to take in the event of a power failure,	
	i.e., use of alternative storage location, backup system or generator.	

### **Evaluation of Training**

Date	Tasks	
Demonstrate how to properly read thermometer and document below (5 occurrences)		
	Temperature:	Verified by:

Upon signing the approval, the trainee and trainer will review the above information and ensure the trainee has demonstrated knowledge and understanding of the above topics.

#### **Approval**

Trainee	Date	
Trainer/ Supervisor	Date	
Comments		



Revision #19, Effective Date 10-20-2025

### **Cannabis Analysis**

This guide is intended to provide a trainee with the necessary skills to perform laboratory analysis of samples suspected to be marijuana, hashish, hashish oil, THC, or THCA-A using a series of examinations. These examinations are stereomicroscopic examination, thin layer chromatography, FTIR, gas chromatography with flame ionization detector and gas chromatography with mass spectral examination.

#### Goals

- To become familiar with the legal status of marijuana in Oklahoma.
- To become skilled at the identification of marijuana samples using a series of examinations.

#### Literature

Date	Literature
	Protocol DR-01 (Cannabis Analysis)
	Protocol DR-01 Attachment 1 – 1% Threshold Testing for Total THC
	Protocol DR-01 Attachment 2 – Flowcharts
	Review Oklahoma Statutes – Title 63, Section 2-101. Specifically, the controlled
	parts of the marijuana plant.
	General familiarization with the remainder of the section.
	Oklahoma Statutes - Title 22, Section 751. Admission of Laboratory findings.
	Release of CDS for independent analysis
	Nakamura, G.R. Forensic Aspects of Cystolithic Hairs of Cannabis and Other
	<b>Plants.</b> Journal of the Association of Official Analytical Chemists. Vol. 52, No. 1,
	1969, pages 5-16
	Mechoulam, R. Marihuana Chemistry. Science. Vol. 168, No. 3936, June 5, 1970,
	pages 1159-1165 (Stop at Biogenesis Section on 3 <sup>rd</sup> page)
	Marihuana, Its Identification. US Treasury Department, Bureau of Narcotics,
	1948
	Methods of Analysis. Internal Revenue Service Publication No. 341, Rev. 6-67,
	page 105



# OSBI Drug Laboratory Training Manual Revision #19, Effective Date 10-20-2025

Lesson Plan #7, Marihuana and THC. Basic Training Program for Forensic Drug
Chemists. May 1972, pages 146-157
Analysis of Drugs. DEA Analytical Manual. U.S. Department of Justice, pages
165-168
Coutts, R.T. & Jones, G.R. A Comparative Analysis of Cannabis Material. Journal
of Forensic Science. Vol. 24, No. 2, 1979, pages 291-302.
Small, E. American Law and the Species Problem in Cannabis. Microgram. Vol.
VII, No. 11, November 1974, pages 131-132.
Nakamura, G.R. and Thornton, J.I. The Forensic Identification of Marihuana:
<b>Some Questions and Answers.</b> <i>Journal of Police Science and Administration.</i> Vol.
I, No. I, 1973, pages 102-112
Zimmerman, Miles C. Marijuana Analysis: Winters V. State. OSBI Legal Update.
Index Tab: Drugs, Control No. 07-77-02, March 24, 1977
Marihuana. Basic Training Program for Forensic Drug Chemists. 2 <sup>nd</sup> Edition, DEA,
pages 6-25 to 6-44
Johnson, Donald W. Hashish Oil. DEA Laboratory Notes. No. 58, May 1973
Cannabis. The Drug Chromatographer. Alltech-Applied Science, Vol. 3, Number
1, 1986, pages 1-3
Recommended Methods for the Identification and Analysis of Cannabis and
Cannabis Products. United Nations Office on Drugs and Crime, 2009
Warner, M.L., Alford, I., Lawrence, D.M., Kohl, A.C., Williams, S.J., Yeatman, D.T.
Comparative Analysis of Freshly Harvested Cannabis Plant Weight and Dried
Cannabis Plant Weight. Forensic Chemistry. Vol. 3, 2017, pages 52-57
Hughes, R. B. M. S. and Kessler, R. R., M.S. Increased Safety and Specificity in
the Thin-Layer Chromatographic Identification of Marihuana. Journal of
Forensic Sciences. Vol. 24, March 19, 1979, pages 842-846
Guy, B. The Identification of Cannabis by Thin Layer Chromatography.
Microgram. Vol. XVII, No. 5, May 1984, pages 78-80



# OSBI Drug Laboratory Training Manual Revision #19, Effective Date 10-20-2025

#### Discussion

Date	Tasks
Microsco	pic Examination
	Microscopic characteristics of marijuana
	Requirement for a positive examination
	Discuss what a typical marijuana leaf looks like, including vein structure
Medical I	Marijuana and Hemp
	Differences between hemp and marijuana
	Are there differences in analysis of medical marijuana and illegal marijuana
	Discuss THC vs THCA
Analysis	
	Proper method for analysis of seeds
	Some of the possible indicators that a second controlled dangerous substance may be present in a marijuana submittal
	How to analyze a green leafy sample
	How to analyze a wax or oil-like substance
	Articulate how to analyze a food/drink product
	The proper procedure for TLC with a sample suspected to contain delta 9 tetrahydrocannabinol
	Articulate interpretation criteria of TLC plate results, such as height of a sample/standard, and color of spots
	When a TLC plate needs to be rejected
	When should a sample use the 1% protocol

### Tasks

Date	Tasks	
Microso	Microscopic Examination	
	Demonstrate use of a stereomicroscope including magnification range	
Analysis	S	
	Demonstrate the preparation and analysis of a sample using TLC	
	Demonstrate how to extract a cannabis sample using BSTFA	
	Prepare TLC Reagent	
	Demonstrate how to use the calibrated pipettes	
	Demonstrate how to prepare a sample for the 1% protocol	



Revision #19, Effective Date 10-20-2025

Approval	
Trainee	Date
Trainer/ Supervisor	Date
Comments	



Revision #19, Effective Date 10-20-2025

### **Drug Literature**

This guide is intended to provide a trainee with the necessary skills to perform laboratory analysis of samples suspected to contain a controlled substance. This analysis will involve the use of various instruments and laboratory techniques to reach conclusions on the identification of a substance.

#### Goals

• To provide a general understanding of controlled substances

#### Literature

Date	Literature
	Review Oklahoma Statutes – Title 63, Section 2-101, 2-201 thru 2-212, 2-321, 2-
	407.1, 2-414, 2-415
	Bell, S. What is a Drug. Forensic Chemistry. pages 213-231, 234-239
	Drugs of Abuse. US Department of Justice, Drug Enforcement Agency, 2011
	Gahlinger, P.M. Illegal Drugs, A Complete Guide to Their History, Chemistry, Use
	and Abuse. pages 232-236 (PDF: Cathinone)
	Synthetic Cathinones ("Bath Salts"). National Institute on Drug Abuse,
	www.drugabuse.gov, 01/10/2013
	Drug Identification Bible. 2022/2023, MDMA, pages 646-652
	Gahlinger, P.M. Illegal Drugs, A Complete Guide to Their History, Chemistry, Use
	and Abuse. pages 264-275 (PDF: Hallucinogens (DMT, Bufotenine and Psilocybin))
	Gahlinger, P.M. Illegal Drugs, A Complete Guide to Their History, Chemistry, Use
	and Abuse. pages 224-228 (PDF: Barbiturates)
	Drug Identification Bible. 2022/2023, Anabolic Steroids, pages 581-584
	Drug Identification Bible. 2022/2023, Heroin, pages 609-623
	Recommended Methods for Testing Opium, Morphine and Heroin. United
	Nations Office on Drugs and Crime, 1998 (Modified PDF version)
	Drug Identification Bible. 2022/2023, Fentanyl, pages 600-605
	Drug Identification Bible. 2022/2023, PCP, pages 653-656
	Drug Identification Bible. 2022/2023, Ketamine, pages 624-626
	Drug Identification Bible. 2022/2023, Amphetamine/Methamphetamine, pages 567-580



Revision #19, Effective Date 10-20-2025

Kelly, B.C. Legally Tripping: A Qualitative Profile of Salvia Divinorum Use Among
Young Adults. Journal of Psychoactive Drugs. Vol. 43 (1), 2011, pages: 46-54
Harris, D. et al. GC-MS Differentiation of Three Synthetic Cannabinoid Positional
Isomers: JWH-250, JWH-302 and JWH-201. Journal of the Clandestine Laboratory
Investigating Chemists Association. Vol. 21, No. 4, October 2011, pages 23-32
Protocol DR-103 (Classification of Synthetic Cannabinoids)

#### Discussion

Date	Tasks
	Convert milligrams to grams
	Vicks inhaler and what substance it contains

Upon signing the approval, the trainee and trainer will review the above information and ensure the trainee has demonstrated knowledge and understanding of the above topics.

#### **Approval**

Trainee	Date	
Trainer/ Supervisor	Date	
Comments		



Revision #19, Effective Date 10-20-2025

### **Pharmaceutical Identification by Literary Reference (DR-5)**

#### Goals

• To establish guidelines when using visual examinations and comparison to a literary reference as the presumptive examinations for tablet and capsule exhibits.

#### Literature

Date	Literature
	Protocol DR-5 (Pharmaceutical Identification by Literary Reference)
	Commonly Abused Prescription Drugs. American Addiction Centers,
	https://drugabuse.com/prescription-drugs/, 05/06/2022
	Commonly Abused Drugs. American Addiction Centers,
	https://drugabuse.com/drugs/most-abused-drugs/, 05/30/2022

#### **Discussion**

Date	Tasks
	When a literary reference is sufficient and when a conclusive analysis is needed
	The procedure when a suspected tablet or capsule was clandestinely
	manufactured
	The different resources that can be used for a literary reference

#### **Tasks**

Look up at least 5 tablet/capsules using the references in DR-05			
	Tablet		Result
Date	Description	Reference	(Identification and Concentration)



**Approval** 

# **OSBI Drug Laboratory Training Manual**

Revision #19, Effective Date 10-20-2025

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Trainee	Date
Trainer/ Supervisor	Date
Comments	



Revision #19, Effective Date 10-20-2025

### **Training Procedures for Extractions & Handling**

#### Goals

• To familiarize the analyst with extraction and handling procedures commonly used to prepare samples for analysis using gas chromatography (GC) and gas chromatographymass spectrometry (GC/MS).

#### Literature

Date	Literature	
	Review SDS sheets on appropriate chemicals used, such Sodium Hydroxide,	
	Hydrochloric Acid, Sodium Bicarbonate, Chloroform, Isopropanol, Hexanes,	
	Methanol, etc.	
	Protocol DR-110 (Extractions) and Appendix C: Extraction Data	
	NIJ Fingerprint Sourcebook, Sections 7.1.5 Evidence Handling and 7.1.6 Packaging	

Date	Tasks
	Which solvent is a good, all-around solvent for most drugs
	Differences in solubility of different drugs
	When other extractions should be used
	<ul> <li>The extraction produces a GC analysis cluttered with peaks</li> </ul>
	<ul> <li>The extraction produces a GC analysis where a secondary peak (i.e., acetaminophen) dwarfs the peak of interest</li> </ul>
	<ul> <li>The extraction produces a GC analysis where poor separation or broad peaks occur</li> </ul>
	<ul> <li>Presumptive color tests indicate amphetamine or methamphetamine</li> </ul>
	<ul> <li>Literary reference indicates a substance that needs to be extracted</li> </ul>
	Why drugs need to be basic extracted
	<ul> <li>Methamphetamine/Amphetamine; Ephedrine/Pseudoephedrine;</li> </ul>
	Phenethylamines
	Which drugs need to be acid extracted
	When to use a back extraction
	Definition of amphoteric and know which drugs exhibit this property
	Which solvent systems should be used with morphine and hydromorphone
	What to do when tablets form an emulsion
	Which substances experience rapid breakdown in solution, and what to do when
	this occurs (i.e., oxymetholone)



# OSBI Drug Laboratory Training Manual Revision #19, Effective Date 10-20-2025

Describe what is to occur if told a sample needs to be placed on the instrument
immediately
Explain how to mix 150 ml of a 2:5:3 solution of A:B:C
Discuss proper packaging for fentanyl and liquids
Discuss working fentanyl on the GC/IRD
What samples are suitable for the GC/IRD
Fentanyl and isomers on the GC/IRD

### Tasks

asks			
Date	Tasks		
Solution	n Preparation		
	Prepare 0.45N NaOH/DI water solution		
	Prepare 10% HCI/DI water solution. H <sub>2</sub> SO <sub>4</sub> can be substituted for HCI		
	Prepare a saturated sodium bicarbonate solution. H <sub>2</sub> SO <sub>4</sub> can be substituted for HCl		
Extraction	ons		
	Determine the pH of a solution		
	Demonstrate the procedure for performing a basic extraction		
	1. Add 0.45N NaOH or another appropriate basic solution to the sample		
	2. Add appropriate amount of chloroform or hexanes		
	3. Mix thoroughly and centrifuge if necessary		
	<ul> <li>Chloroform is preferred over hexanes since it can solubilize a greater</li> </ul>		
	number of drugs		
	4. When utilizing this extraction procedure chloroform will form the bottom		
	layer and hexanes will form the top layer		
	Demonstrate the procedure for performing an acidic extraction		
	1. Add 5-10 drops of 10% HCl or H₂SO₄ to the sample		
	2. Add appropriate amount of chloroform		
	Mix thoroughly and centrifuge if necessary		
	Demonstrate the procedure for a back extraction		
	1. Add 1 milliliter of DI water to a sample		
	2. Add 5-10 drops of 10% HCl solution to the sample		
	3. Mix well and centrifuge if necessary		
	4. Remove the aqueous layer and place in another culture tube		
	5. Make the aqueous layer basic by adding 0.45N NaOH solution		
	6. Verify the pH of the aqueous solution has converted from an acid to a base		
	7. Once basic, add an appropriate solvent (chloroform or hexanes)		
	8. Mix well and centrifuge if necessary		



Revision #19, Effective Date 10-20-2025

Date	Demonstrate		
Sample	Sample Handling		
	Demonstrate how to label evidence and test tubes/vials		
	Demonstrate how to dilute and concentrate a sample		
	Safely remove clean syringe with needle (previously prepared by trainer) from		
	sharps container, rinse with a solvent, recap and replace back into container		
	Safely remove razor or knife from sharps container (previously prepared by		
	trainer), sample (i.e., with swab) and replace back into container.		
	Safely remove broken glass from envelope (previously prepared by trainer), sample		
	residue from broken glass, and place into an appropriate sharps container		
	Demonstrate handling, marking, & sampling of evidence to be forwarded to Latent		
	Evidence Unit, with minimal risk of damaging potential latent prints		

Approval	
Trainee	Date
Trainer/ Supervisor	Date
Comments	



Revision #19, Effective Date 10-20-2025

### **Identification of Gamma-Hydroxybutyric Acid through Derivatization (DR-7)**

#### Goals

• To establish guidelines for the identification of Gamma-Hydroxybutyric Acid (GHB) through derivatization with bis-(trimethylsilyl) trifluoroacetamide (BSTFA) and analysis using gas chromatography (GC) and gas chromatography mass spectrometry (GC/MS).

#### Literature

Date	Literature	
	Protocol DR-7 (Identification of Gamma-Hydroxybutyric Acid through	
	Derivatization)	
	SDS sheets for BSTFA and acetonitrile	
	Drug Identification Bible. 2022/2023, GHB, pages 606-608	
	Kilpatrick, G. A. GHB. State Police-San Francisco.	
	Bell, S. Derivatization. Forensic Chemistry. pages 203-205	
	Bommarito, C. Analytical Profile of Gamma-Hydroxybutyric Acid (GHB).	
	Journal of the Clandestine Laboratory Investigating Chemists Association. Vol. 3,	
	No. 3, July 1993, pages 10-12	
	Pearson, J.R., Reid, E.F., & Rowe, J.E. The Preparation of Y-Butyrolactone from	
	Readily Available Starting Materials. Journal of the Clandestine Laboratory	
	Investigating Chemists Association. Vol. 19, No. 1, January 2009, pages 8-13	

Date	Tasks
	The theory of derivatization and why it is performed in the differentiation of GHB and GBL
	How to recognize when GHB may be present in a sample
	The reason for washing a liquid sample suspected of containing GHB and/or GBL with chloroform
	How heat or pH may affect a sample containing GHB and/or GBL
	Proper reporting results for both dry and liquid samples



Revision #19, Effective Date 10-20-2025

#### **Tasks**

Date	Tasks (If samples are available)
	Demonstrate derivatization procedure on a dry sample
	Demonstrate derivatization procedure on a liquid sample
	Demonstrate ability to identify GBL if present in a liquid sample

Approval	
Trainee	Date
Trainer/ Supervisor	Date
Comments	



Revision #19, Effective Date 10-20-2025

### **Training Procedures for Color Tests (DR-10, DR-11, DR-13)**

#### Goals

• To establish guidelines for preliminary screening tests that respond to particular functional groups on substances causing characteristic color changes. These examinations can give the analyst a basis as to which extractions and/or examinations are necessary for further conclusive instrumental analysis.

#### Literature

Date	Literature
	Protocol DR-10 (Color Tests: Marquis)
	Protocol DR-11 (Color Tests: Cobalt Thiocyanate)
	Protocol DR-13 (Color Tests: Bates)
	Clarke's Analysis of Drugs and Poisons, Third Edition, pages 279 -300

Date	Task
	The specificity of color tests and their role in drug analysis
	Where all color tests are to be performed
	Reasons why a color test may be negative for a compound even when the
	compound is present in a sample
	Indications of when a reagent may need to be discarded and new reagent made
	The procedure for performing a negative control and when it is necessary
	The procedure for performing a positive control and when it is necessary



Revision #19, Effective Date 10-20-2025

#### **Tasks**

	1		
Tasks			
		Cobalt	
	Marquis Test	Thiocyanate	Bates' Test
Demonstrate the			
procedure for			
making the reagent			
Demonstrate the			
quality control			
verification and			
documentation			
procedure			
Demonstrate the			
procedure for			
performing the test			
Based on the			
observed results of			
the color test,			
articulate a suitable			
extraction			
procedure			

Upon signing the approval, the trainee and trainer will review the above information and ensure the trainee has demonstrated knowledge and understanding of the above topics.

### **Approval**

Trainee	Date
Trainer/ Supervisor	Date
Comments	



Revision #19, Effective Date 10-20-2025

### **Cocaine Free-base Determination by Hexane Solubility (DR-21)**

#### Goals

- To establish a knowledge of cocaine and cocaine base.
- To establish guidelines to differentiate cocaine hydrochloride from cocaine free base based on their solubility in hexanes.

#### Literature

Date	Literature
	Protocol DR-21 (Cocaine Free Base Determination by Hexane Solubility)
	Gahlinger, P.M. Illegal Drugs, A Complete Guide to Their History, Chemistry, Use
	and Abuse. pages 240-254 (PDF: Cocaine)
	Cocaine. NIDA InfoFacts. www.drugabuse.gov, 01/10/2013
	Recommended Methods for Identification and Analysis of Cocaine in Seized
	Materials. United Nations Office on Drugs and Crime, 2012
	Crack Cocaine Recipe, www.hyperreal.org, 1992
	Drug Identification Bible. 2022/2023, Cocaine, pages 585-599

Date	Tasks
	Why cocaine hydrochloride and cocaine base need to be differentiated
	What color tests can be used to indicate the presence of cocaine or cocaine base
	The procedure if a sample is negative for cocaine base
	The difference between cocaine base and cocaine HCl



Revision #19, Effective Date 10-20-2025

#### **Tasks**

Date	Tasks
	Demonstrate the difference in solubility of cocaine hydrochloride and cocaine base
	using both methanol and hexanes
	Demonstrate the difference in reactivity of cocaine hydrochloride and cocaine base
	with the Cobalt Thiocyanate color test and Bates Test

Approval	
Trainee	Date
Trainer/ Supervisor	Date
Comments	



Revision #19, Effective Date 10-20-2025

### **Identification of Lysergic Acid Diethylamide (DR-101)**

#### Goals

- To establish a knowledge of Lysergic Acid Diethylamide (LSD).
- To familiarize the analyst with procedures used in the identification of LSD in different forms.

#### Literature

Date	Literature	
	Protocol DR-101 (Identification of Lysergic Acid Diethylamide)	
	Drug Identification Bible. 2022/2023, LSD, pages 627-630	
	Recommended Methods for Testing Lysergide (LSD). United Nations Office on	
	Drugs and Crime, 1989 (Modified PDF version)	
	Smith, F., Handbook for Drug Analysis. 2005, pages 186-187	

#### Discussion

Date	Tasks	
	Presumptive test for LSD on a sample	
	Which GC/MS methods would be used for an LSD sample and reagent blank	
	Explain how to sample blotter paper	
	What report wording is used when analyzing suspected LSD	

#### **Tasks**

Date	Tasks
	Analyze LSD, LAMPA and a mixture of LSD & LAMPA to demonstrate the differences
	of the compounds on the GC and GC/MS

Upon signing the approval, the trainee and trainer will review the above information and ensure the trainee has demonstrated knowledge and understanding of the above topics.

#### **Approval**

Trainee	Date
Trainer/ Supervisor	Date
Comments	



Revision #19, Effective Date 10-20-2025

### **Gas Chromatography Analysis (Flame Ionizing Detector) (DR-30)**

#### Goals

- To gain knowledge of the theory of gas chromatography and how to use it as a non-confirmatory test in the analysis of submitted samples.
- To establish guidelines for the gas chromatograph maintenance.
- To demonstrate a gas chromatograph is working properly by using quality assurance and quality control methods.
- The analyst will learn how to interpret data from this type of analysis.

#### Literature

Date	Literature
	Protocol DR-30 (Gas Chromatography Analysis (flame ionization detector))
	Basic Operation of the Gas Chromatograph (Appendix I of Training Manual)
	Bell, S. Forensic Chemistry. pages 192-200 (PDF: GC and MS)

Date	Tasks	
Theory of Gas Chromatography		
	The injection port	
	Expansion volumes and how to determine the appropriate injection volume	
	Megabore columns used in the gas chromatograph (DB-1 and DB-50)	
	The flame ionizing detector	
	Split ratios and why it is used with the instrument	
	The different methods used for analysis, such as Drug1, Extend1 and Method1	
	and/or any other methods being used for analysis	
	Make-up gas	
	Retention time and how it applies to gas chromatograph analysis	
Controls		
	When standard ladders are to be run	
	What is the negative control and when should it be ran	
	What is the positive control and when should it be ran	
	When methanol blanks should be run	
	The proper corrective procedure if the retention time of the cocaine standard	
	exceeds plus or minus 2% of the standard ladder	
	What, if any, type of extraneous peaks is allowed in cocaine standard	
	What constitutes contamination/carryover in methanol blanks	
	What are the acceptable levels or sizes of contamination/carryover peaks allowed	



# OSBI Drug Laboratory Training Manual Revision #19, Effective Date 10-20-2025

	The proper corrective procedure if contamination occurs during methanol blank
	runs
	The proper corrective procedure if the instrument fails to produce a satisfactory
	chromatogram for the cocaine standard
Misc.	
	What is the temperature programming and where can it be found
	When must a standard be run on the gas chromatograph
	The maximum number of days between running a standard and a sample on the
	gas chromatograph
	When changing of the liner and septa is required
	When changing of the gold seal is required
	When cleaning of the flame ionization detector is required
	When cleaning of the injection port is required
	What conditions and/or situations destroy the stationary phase of a GC column
	Articulate reasons that GC data may be "rejected"
	What happens to sample after analysis through instrument?
	<ul> <li>Where does the waste from split vent go?</li> </ul>
	<ul> <li>Where does sample go after leaving the jet?</li> </ul>

### Tasks

Date	Tasks	
Maintenance		
	Change the liner and septum and reset macro counts	
	Change the gold seal, clean the injector port and septa nut and reset macro counts	
	Clean the flame ionization detector	
	Change a column when necessary	
	Change the split vent filter and line	
	Demonstrate syringe replacement	
	Demonstrate proper wash and waste bottle volumes	
	Extract standard ladder, run on GC and update macros	
	Record any maintenance performed on the proper maintenance log	
Use of t	he Instrument	
	Properly prepare a sample in an auto-sampler vial	
	Properly load a sample for analysis on the gas chromatograph, including entering	
	information in sequence log	
	Demonstrate the proper use of controls:	
	MeOH blank	
	Cocaine standard	
	Reagent blank	



Revision #19, Effective Date 10-20-2025

Spectral Interpretation		
	The analyst will review instrumental data and discuss with the trainer what is and is	
	not acceptable for casework analysis	
	<ul> <li>Samples outside of the 2% retention time window</li> </ul>	
	<ul> <li>Demonstrate calculating the 2% window based on the retention time of a</li> </ul>	
	standard	
	Peak separation	
	Acceptable/unacceptable chromatography	
	How to integrate a peak in data analysis	
	* See Training folder for examples of instrumental data that might need more work	
	\\VM-FSC-FILES\Common\DrugLab\1 Technical Manager\1 Training	
	Properly interpret the data obtained and explain how to apply this data if further	
	analysis is required	
	Demonstrate the proper reporting of results available from gas chromatographic	
	data	

Approval		
Trainee	Date	
Trainer/ Supervisor	Date	
Comments		



Revision #19, Effective Date 10-20-2025

# Analysis of Mushrooms to Determine the Presence of Psilocyn or Psilocybin (DR-45)

#### Goals

• To establish guidelines for the differentiation of psilocyn and psilocybin for identification.

#### Literature

Date	Literature
	Protocol DR-45 (Analysis of Mushrooms to Determine Presence of Psilocyn or
	Psilocybin)
	Drug Identification Bible. 2022/2023, Peyote & Psilocybin Mushrooms, pages 657-
	663
	Recommended Methods for Testing Peyote Cactus (Mescal Buttons)/Mescaline
	and Psilocybe Mushrooms/Psilocybin. United Nations Office on Drugs and Crime,
	1989 (Modified PDF version)

Date	Tasks	
	Why psilocyn and psilocybin need to be differentiated	
	What happens to psilocyn and psilocybin when injected into the gas	
	chromatograph in methanol	
	The different methods that can be used to differentiate psilocyn and psilocybin	
Thin Lay	Thin Layer Chromatography Examination	
	Articulate proper procedure and specificity for TLC with a sample suspected to	
	contain psilocyn or psilocybin	
	Articulate familiarization with Rf value	
	Articulate other procedure(s) that has to be performed in conjunction with TLC	
	with a sample suspected to contain psilocyn or psilocybin	
Derivat	Derivatization of Sample	
	Articulate proper procedure and specificity for derivatization with a sample	
	suspected to contain psilocyn or psilocybin	



Revision #19, Effective Date 10-20-2025

#### **Tasks**

Approval

Date	Tasks	
Thin La	Thin Layer Chromatography Examination	
	Prepare Mushroom TLC Reagent	
	Demonstrate the preparation and analysis of a sample using TLC	
	Demonstrate the two ways for visualization of a TLC plate for a sample suspected	
	to contain psilocyn and/or psilocybin	
	Establish interpretation criteria of TLC test results, such as height of a	
	sample/standard, and color of spots	
Derivat	Derivatization of Sample	
	Demonstrate derivatization of a sample suspected to contain psilocyn and/or	
	psilocybin	

Trainee	Date	
Trainer/ Supervisor	Date	
Comments		



Revision #19, Effective Date 10-20-2025

### PDF Examination Documentation Procedure (DR-50)

#### Goals

- To establish guidelines for generating, storing, transferring and attaching instrumental data into the BEAST Image Vault.
- The analyst will learn the security and tracking features associated with this process.

#### Literature

Date	Tasks
	Protocol DR-50 (PDF Examination Documentation Procedure)
	Review DRQM-11 Examination Documentation

#### Discussion

Date	Tasks
	Articulate the security and tracking features associated with this process (creation,
	merging, and uploading of PDFs)
	What the validation code is and where it comes from
	Articulate the manner for archiving PDFs

#### **Tasks**

Date	Tasks	
	Set up PDF folders on the instrument computer  Establish a method of transferring PDFs from the instrument computer to the analyst's computer	
	Demonstrate transferring PDF files from the instrument computer to the analyst's computer	
	Acquire PDF editing software  Demonstrate the merging of PDFs  Demonstrate the naming of PDFs  Demonstrate how to make a correction to a PDF	
	Demonstrate uploading a PDF file into the BEAST Image Vault	
	Demonstrate removing a PDF file from the BEAST Image Vault	



**Approval** 

# **OSBI Drug Laboratory Training Manual**

Revision #19, Effective Date 10-20-2025

Trainee	Date
Trainer/ Supervisor	Date
Comments	



Revision #19, Effective Date 10-20-2025

## **Drug Analysis by FTIR (DR-60)**

#### Goals

- To learn the theory of FTIR and how to use it as a confirmatory test in the analysis of submitted samples.
- To familiarize the analyst with how to maintain the FTIR instrument and ensure that it is working properly by using quality assurance and quality control methods.
- The analyst will learn how to interpret data from this type of analysis.

#### Literature

Date	
	Literature
	Protocol DR-60 (Drug Analysis by FTIR)
	Thermo Scientific. FT-IR Glossary. (PDF)
	Thermo Scientific. Introduction to Fourier Transform Infrared Spectroscopy (PDF)
	LCGC ChromAcademy. Introduction to Infrared Spectroscopy. (PDF)
Perkin Elmer, FT-IR Spectroscopy Attenuated Total Reflectance (ATR). (PDF Bell, S. Spectroscopy. Forensic Chemistry. pages 149-159	Perkin Elmer, FT-IR Spectroscopy Attenuated Total Reflectance (ATR). (PDF)
	Bell, S. <b>Spectroscopy.</b> Forensic Chemistry. pages 149-159
	Bell, S. Infrared Spectroscopy. Forensic Chemistry. pages 161-169
	Hugel, J., Meyers, J.A. & Lankin, D.C. Analysis of the Hallucinogens, Infrared (IR)
	Spectroscopy. Handbook of Forensic Drug Analysis. 2005, pages 154-164
	Clarke's Isolation and Identification of Drugs. Vol. I, 3rd Edition, pages 328-344

#### Discussion

Date	Tasks
Theory	of FTIR
	Theory and definition of FTIR
	Define:
	Wavelength
	Wavenumber
	Interferometer
	Constructive Interference (pertaining to FTIR)
	Destructive Interference (pertaining to FTIR)



	Explain how the following pertain to FTIR
	• Gain
	Resolution
	Single beam spectrum
	Sensitivity
	Describe the components of the FTIR and their function
	The differences between transmission and reflectance modes
	What wavelength range is analyzed using FTIR
	Explain evanescent wave as it pertains to FTIR ATR
	How the interferometer functions
	Describe what a laser is and its function in FTIR, including its travel path within the FTIR
	What is a background and why is it collected
	Describe what ATR is
	<ul> <li>How the ATR functions including:</li> <li>What the crystal is made of and why it doesn't interfere with analysis</li> <li>Why CO<sub>2</sub> and H<sub>2</sub>O can still appear in a spectrum after background scans have been collected</li> </ul>
	Describe what the FTIR does to the sample during analysis
	Why FTIR on tablets can be difficult
Controls	
	When spectra of a polystyrene standard will be obtained and what will it be compared to
	The proper corrective procedure when the polystyrene standard fails
	When a background spectrum will be collected
	List ways contamination may be identified on the stage crystal and the tower arm
	What steps are taken to ensure the ATR is free from contamination
	Describe the spectra of a "blank"



Misc.	
	Describe how to prepare a sample for FTIR analysis using the ATR
	Explain how humidity can affect the spectra and the quality of the match
	Explain how to remove moisture from a sample
	What must be done if analysis by FTIR does not indicate a controlled dangerous substance
	Articulate why a second sample would be taken and what documentation is needed to be included in the casefile
	Articulate reasons that FTIR data may be "rejected"

## Tasks

Date	Tasks	
Mainte	enance	
	Demonstrate the analysis of a polystyrene standard which is to be analyzed using the "OSBI Macro," before casework is performed.	
	Demonstrate proper cleaning of the ATR	
	Demonstrate checking the humidity levels inside the instrument	
Use of	Jse of the Instrument	
	Demonstrate preparation of a sample for analysis	
	Demonstrate collection of background scans	
	Demonstrate collection of scans using the ATR accessory	
Spectra	al Interpretation	
	Review instrumental data and discuss what is and is not acceptable for casework analysis	
	Determine if CO <sub>2</sub> and/ or H <sub>2</sub> O are present in the scans	
	Determine if another compound is present in the scans	
	Demonstrate proper report writing	



Revision #19, Effective Date 10-20-2025

Upon signing the approval, the trainee and trainer will review the above information and ensure the trainee has demonstrated knowledge and understanding of the above topics. **Approval** 

Trainee	Date
Trainer/ Supervisor	Date
Comments	



# **Requirements Prior to Drug Analysis Using FTIR**

## **Sample Analysis**

Date	Tasks
	Analyze at least 30 practice samples and record all results;
	document using an Excel spreadsheet and archive in analyst's folder on QA server

## **Evaluation of Training**

Date	Tasks	
	Complete and review a competency test, with accurate results	
	Complete a technical questions session with a minimum score of 80%, with	
	Technical Manager or Appointee	
	Average Score	

Approval		
Trainee	Date	
Trainer/ Supervisor	Date	
Comments		



Revision #19, Effective Date 10-20-2025

## **Gas Chromatograph Mass Spectrometer Methods for Drug Analysis (DR-70)**

#### Goals

- To learn the theory of gas chromatography mass spectrometer and how to use it as a confirmatory test in the analysis of submitted samples.
- To familiarize the analyst with how to maintain the gas chromatograph mass spectrometer instrument and ensure that it is working properly by using quality assurance and quality control methods.
- The trainee will learn how to interpret data from this type of analysis.

\*\*Prior to beginning, the trainee must complete GC training (DR-30)\*\*

#### Literature

Date	Literature
	Protocol DR-70 (Gas Chromatograph Mass Spectrometer Methods for Drug Analysis)
	Basic Operation of the Mass Spectrometer (Appendix II of Training Manual)
	Isomers (Appendix V of Training Manual)
	Hugel, J., Meyers, J.A. & Lankin, D.C. Analysis of the Hallucinogens, Mass
	Spectrometry. Handbook of Forensic Drug Analysis. 2005, pages 176-185
	Pavia, D.L., Lampman, G.M., Kriz, G.S. Mass Spectrometry. Intro to Spectroscopy.
	2001, pages 390-398 and 446-448
	Optional: Prall, J. D. & Cardone. Gas Chromatography/Mass Spectrometric (GC/MS)
	Analysis of Drugs Using Spectral and Retention Index Matching. DEA Central Lab,
	Dallas, TX and FAA Toxicology and Research Lab, OKC, OK.
	(No date and reference available)

#### Discussion

Date	Tasks
Theory	of Mass Spectrometry
	The different methods and when to use for analysis, i.e., drug100, drug200, LSD,
	extnd100, low100, meth100, and/or any other methods being used for analysis
	Split ratio and why it is used with the instrument
	Capillary columns used in the mass spectrometer
	Nonpolar and polar liquid phases
	How does film thickness affect the performance
	Splitting of the molecule and the function of the ion source
	The quadrupole mass filter and how it functions
	The electron multiplier



	Retention time and retention index, and how it applies to the GC/MS	
	What is the hydrocarbon ladder and what role does it play in the retention index	
	How the retention index is calculated	
	The maximum allowed difference of the calculated Retention Index Difference	
011	What carrier gas does the Drug Lab use	
Controls		
	What is the negative control and when should it be ran	
	What is the positive control and when should it be ran	
	When solvent blanks are to be run	
	The proper corrective procedure if the retention index of the Cocaine standard exceeds plus or minus 2%	
	The proper corrective procedure if contamination occurs during the daily solvent blank run	
	The proper corrective procedure if contamination occurs during a casework solvent blank run	
	When the Tune Eval/Autotune are to be run	
	The proper corrective procedure if erroneous assignment of mass values to fragments in a sample, standard, or autotune	
	The proper corrective procedure for the failure to produce a satisfactory cocaine spectrum for the cocaine standard	
	The proper corrective procedure if contamination peaks are found on the autotune (m/e 18, 44, etc.)	
	When running the hydrocarbon ladder is required	
	When changing the liner and septum are required	
	When changing the gold seal is required and the difference between the GC and the GC/MS	
Misc.	·	
	What is the temperature programming and where can it be found	
	The theory of a mass spectral library and where the libraries come from	
	When does a standard have to be run	
	What information has to be retained with a new standard	
	What information about the standard is listed on each mass spectra printout	
	Articulate reasons that data from the GC/MS may be "rejected"	
	What happens to sample after analysis through instrument?	
	<ul> <li>Where does the waste from split vent go?</li> </ul>	
	<ul> <li>Where does sample go after leaving the mass spec?</li> </ul>	



Revision #19, Effective Date 10-20-2025

Define	:
•	Structural Isomers

- Stereoisomers
- Enantiomers/Optical Isomers
- Diastereomers/Geometric Isomers

### **Tasks**

Date	Tasks
Mainte	nance
	Demonstrate how to record maintenance performed in the instrument
	maintenance log
	Demonstrate how to prepare and run the hydrocarbon ladder and ensure the
	appropriate retention times have been updated in the macro
	Demonstrate changing liner and septum, resetting macro counts
	Demonstrate changing gold seal, cleaning injection port and septa nut, and resetting macro count
	Demonstrate how to autotune instrument, interpret the data and save the PDFs
	Demonstrate how to vent the mass spec
	Demonstrate how to disassemble and clean ion source, replace filaments, and
	reassemble
	Demonstrate how to pump down the mass spec
	Demonstrate how to change the split vent filter and line
	Demonstrate how to change a column
Use of t	he Instrument
	Demonstrate how to properly dilute or concentrate a sample for GC/MS analysis
	Demonstrate how to properly load a sample for analysis on the GC/MS, including
	entering information in the Sequence log
	Demonstrate the proper use of controls:
	Reagent Blanks
	Cocaine Standards
	Articulate the requirements for Reagent Blanks and the Cocaine Standard
	What constitutes an acceptable Blank?
	What constitutes an acceptable Cocaine Standard?
	What methods are used and when/why?
	Demonstrate the interpretation and comparison of the mass spectral data received
	from the instrument



Revision #19, Effective Date 10-20-2025

Demonstrate how to perform a background subtraction and articulate when a background subtraction is needed.
Demonstrate how to perform a manual scan. Articulate when a manual scan may be needed and what documentation is required if a manual scan is performed.
Demonstrate the proper reporting of results from the data received from the instrument

Spectral	Interpretation	
	The trainee will review instrumental data and discuss with the trainer what is and is	
	not acceptable for casework analysis	
	Peaks past the molecular ion peak	
	Background subtraction	
	Complete spectra	
	Extra/absent ions in a spectrum	
	Calculate retention index difference	
	* See Training folder for examples of instrumental data that might need more work	
	\\VM-FSC-FILES\Common\DrugLab\1 Technical Manager\1 Training	
	What are possible sources of high background and how do you fix it	
	Why would you experience broad, misshaped, or tailing peaks and how do you fix	
	each one	

Upon signing the approval, the trainee and trainer will review the above information and ensure the trainee has demonstrated knowledge and understanding of the above topics.

## **Approval**

Trainee	Date
Trainer/ Supervisor	Date
Comments	



Revision #19, Effective Date 10-20-2025

## **Administrative and Technical Reviewing of Casework**

#### Goals:

• To provide the trainee with knowledge and skills necessary to perform Administrative and Technical Reviews on another Analyst's casework

#### Literature

Date	Literature
	Review OSBI QP 31, Reviews
	Review DRQM-12 Case Reviews

#### Discussion

Date	Tasks
	Discuss with Trainer how to perform a Technical Review of a Drug Case
	Discuss how different prosecutorial charges affect the identification of certain
	compounds (i.e., pseudoephedrine vs. ephedrine, trafficking, distribution, etc.)

#### Tasks

Date	Tasks		
Observe t	Observe three Analysts perform Admin/Tech Reviews (5 cases each Analyst)		
	Name of Analyst 1:		
	Name of Analyst 2:		
	Name of Analyst 3:		

Upon signing the approval, the trainee and trainer will review the above information and ensure the trainee has demonstrated knowledge and understanding of the above topics.

Approval		
Trainee _	 Date	
Trainer/ Supervisor _	Date	
Comments _		



## **Requirements Prior to Drug Analysis**

## **Sample Analysis**

Date	Tasks
	Perform the visual examination on approximately 25 green leafy samples that
	includes both negatives and positives, analyze using thin layer chromatography,
	and record all results; document using an Excel spreadsheet and archive in the
	analyst's folder on QA server.
	All results must be accurate; if not then documentation of trainee & trainer review
	must be completed with explanation of possible differences & TM notified.
	Complete a competency for the calibrated pipettes. A satisfactory competency test
	will be within 10% of the expected value. The Technical Manager or trainer will
	prepare the competency and record the expected value.
	Analyze approximately 100 provided samples on the GC and GC/MS, to include
	positive and negative controlled dangerous substances and record all results;
	document using an Excel spreadsheet and archive in the analyst's folder on QA
	server.
	All results must be accurate; if not then documentation of trainee & trainer review
	must be completed with explanation of possible differences & TM notified.
	Analyze approximately 70 practice drug cases, on the GC and GC/MS, and record all
	results; document using an Excel spreadsheet and archive in the analyst's folder on
	QA server.
	All results must be accurate; if not then documentation of trainee & trainer review
	must be completed with explanation of possible differences & TM notified.
	Perform a training technical review on 50 cases.
	These cases will be routed to the trainee in the BEAST using the route for training
	purposes code. The technical review will be documented on an Excel spreadsheet.
	The reviews done by the trainee will be compared to the official technical review
	by the TM or the Trainer.

## **Evaluation of Training**

Date	Tasks	
	Complete and review a competency test, with accurate results	
	Complete a technical questions session with a minimum score of 80%, with	
Technical Manager or Appointee		
	Average Score	
	Complete a mock trial session, with approval from Technical Manager	



Approval	
Trainee	Date
Trainer/ Supervisor	Date
Comments	



Revision #19, Effective Date 10-20-2025

## **Liquid Nitrogen Safety**

Liquid Nitrogen is the liquefied form of nitrogen gas. When nitrogen is in the gas phase, it is mostly inert gas that is colorless, odorless, and tasteless. In the liquid phase, nitrogen is very cold (-196 C or -320 F), which makes it ideal for keeping things cool. Because of its extremely cold temperature, any exposure to your skin can cause severe frostbite. On vaporization, liquid nitrogen expands by a factor of almost 700, so 1 liter of liquid nitrogen becomes 24.6 cubic feet of nitrogen gas. This can cause explosion of a sealed container, or it can displace oxygen in the room and cause suffocation without warning. Liquid nitrogen should always be stored in a vented container in a well-ventilated room. Oxygen may condense on the surface of liquid nitrogen causing it to be highly reactive with organic materials. This can cause ordinarily noncombustible materials to burn rapidly when it comes in contact with oxygen enriched liquid nitrogen. When handling liquid nitrogen, always wear thermal gloves and a protective face shield. Never dispose of liquid nitrogen by pouring it on the floor as it could displace enough oxygen to cause suffocation. Nitrogen gas is colorless and odorless, the cloud that forms when liquid nitrogen is poured on the floor is condensed water vapor from the air, not nitrogen gas.

#### Goals

- To become knowledgeable of the hazards of liquid nitrogen
- To learn the safety precautions to utilize when handling liquid nitrogen
- To learn how to properly use/transfer liquid nitrogen

#### Literature

Date	Literature
	Safety Data Sheet for liquid nitrogen
	OSBI Policy 121.1 Appendix I, personal protective equipment required when
	handling liquid nitrogen

#### Discussion

Date	Tasks	
Why is liquid nitrogen used in the laboratory		
	What are three hazards of liquid nitrogen	
	What personal protective equipment must be used when handling liquid nitrogen	
· · · · · · · · · · · · · · · · · · ·	What is the proper type of container to use to transport liquid nitrogen	
	What first aid is necessary if liquid nitrogen spills on skin or eyes	
How do you report a liquid nitrogen injury if it occurs		
	Why is there a safety release valve on the cryogenic cylinder	
	Why must you never use a tight-fitting cap on a dewar of liquid nitrogen	



Revision #19, Effective Date 10-20-2025

### **Tasks**

Date	Tasks
	Trainer will demonstrate the proper technique for transferring liquid nitrogen from
	the cryogenic cylinder to a small dewar.

## **Evaluation of Training**

Demonstrate how to properly transfer liquid nitrogen	
--	--

Upon signing the approval, the trainee and trainer will review the above information and ensure the trainee has demonstrated knowledge and understanding of the above topics.

### **Approval**

Trainee	Date
Trainer/ Supervisor	Date
Comments	



Revision #19, Effective Date 10-20-2025

## **Gas Chromatograph Infrared Detector Methods for Drug Analysis (DR-75)**

#### Goals

- To learn the theory of gas chromatography infrared detector and how to use it as a confirmatory test in the analysis of submitted samples.
- To familiarize the analyst with how to maintain the gas chromatograph infrared detector instrument and ensure that it is working properly by using quality assurance and quality control methods.
- The trainee will learn how to interpret data from this type of analysis.

#### Literature

Date	Literature
	Protocol DR-75 (Gas Chromatograph Infrared Detector Methods for Drug Analysis)
	IRD3 Operations Manual, Rev 1-3, ASIC
	Essential FTIR Operations Manual for GC IRD Users, Rev 0-3
	IRD 3 Hardware & Schematic Overview

#### **Tasks**

Date	Tasks		
Mainte	nance		
	Demonstrate how to fill the GCIRD with liquid nitrogen		
Use of	the Instrument		
	Demonstrate how to properly load a sample for analysis on the GCIRD, including		
	entering information in the Sample Table		
	Demonstrate the proper use of controls:		
	Solvent Blanks		
	Cocaine Standards		
	Demonstrate the interpretation and comparison of data		

<sup>\*\*</sup>Prior to beginning, the trainee must complete GC (DR-30), FTIR training (DR-60), and Liquid

Nitrogen Safety\*\*



Revision #19, Effective Date 10-20-2025

## Requirements Prior to Drug Analysis using the GCIRD

## **Sample Analysis**

Date	Tasks
Analyze at least 30 practice samples and record all results;	
	document using an Excel spreadsheet and archive in analyst's folder on QA server

### **Evaluation of Training**

Date	Tasks
	Complete and review a competency test, with accurate results

Upon signing the approval, the trainee and trainer will review the above information and ensure the trainee has demonstrated knowledge and understanding of the above topics.

Approval	
Trainee	Date
Trainer/ Supervisor	Date
Comments	



Revision #19, Effective Date 10-20-2025

### **Controlled Substances Technician**

The Controlled Substances Technician's job is to assist the analysts and supervisors in the Controlled Substances Unit. This may include a variety of duties, including but not limited to: making reagents, checking scales & refrigerator temperatures, instrument maintenance and preparation of sampling apparatuses. Education and Experience Requirements: At a minimum, the Controlled Substances Technician is required to have graduated from high school or have an equivalent diploma. It is preferred the Technician has attended or is currently attending an accredited college or university, with preferred coursework in chemistry, forensic science, criminalistics, toxicology or a closely related natural science.

The Controlled Substances Technician will complete portions of the same sections of this training manual as a Criminalist in the Controlled Substances Unit. The trainee will date when each assignment is completed. The trainee and trainer will sign/date the Approvals and initial/date the Checklist when the required sections have been completed. Required sections will be listed in the Checklist.

The trainer will demonstrate to the Controlled Substances Technician some of the tasks that will be required duties; there may be no associated section in the training manual and no reading associated with those tasks. By signing off on the Checklist, the trainee has demonstrated their ability of completing the task to the trainer.

A competency test must be completed, with anticipated results, prior to being released to create items that could be used for testing. The competency may include testing performed by an authorized analyst of the item(s) used for testing.

The Technician may be authorized to perform some duties before all sections of the Checklist are completed.

	Trainee	Trainer	
Section	Initials	Initials	Date
Weights and Measures Utilized			
Read DR-3			
Read DR-4 (DR-4 Attachments not required)			
Demonstrate how to properly use & document weights:			
Bench Scale			
Large Capacity Scale			
Analytical Scale			
Complete a 31-day Measurement Study on the three scales			



Thermometers	
All sections	
Extractions & Handling	
Literature	
Solution Preparation of 0.45 N Sodium Hydroxide	
Solution Preparation of Concentrated Sodium Bicarbonate	
Solution Preparation of Concentrated Sodium Hydroxide	
Determination of pH of solution	
Demonstrate the procedure for basic extraction	
Color Tests	
Read DR-10	
Read DR-11	
Read DR-13	
Solution Preparation of Marguis Reagent	
Demonstrate the quality control verification and	
documentation for Marquis Reagent	
Solution Preparation of Cobalt Reagent	
Demonstrate the quality control verification and	
documentation for Cobalt Reagent	
Demonstrate the quality control verification and	
documentation for Bates Test	
Gas Chromatography	
Read DR-30	
Maintenance Section of Training Manual	
Analysis of Mushrooms	
Solution Preparation of Mushroom TLC Reagent	
Solution Preparation of Ehrlichs	
Demonstrate the verification of the TLC Reagent and	
Ehrlichs	
Drug Analysis by FTIR	
Read DR-60	
Maintenance Section of Training Manual	
Gas Chromatograph Mass Spectrometer	
Read DR-70	
Maintenance Section of Training Manual	
Miscellaneous	
Recycling (shredded paper, cardboard, etc.)	
Refill the hydrogen generator bottles	
Check temperatures on the refrigerators	



Revision #19, Effective Date 10-20-2025

Gases
Checking Eyewashes and documenting
Printing reports
Checking Safety Showers and documenting
Decontaminating phones and documenting

Upon signing the approval, the trainee and trainer will review the above information and ensure the trainee has demonstrated knowledge and understanding of the above topics.

Approval	
Trainee	Date
Trainer/ Supervisor	Date
Comments	



Revision #19, Effective Date 10-20-2025

### **Appendix I - Basic Operation of the Gas Chromatograph**

In order to achieve maximum resolution between similar compounds on the gas chromatograph (GC), basic understanding of certain variables should be understood. Clear separation must be obtained for the mass spectrometer (MS) to distinguish between a mixture of molecular compounds and a single molecular compound. The capillary gas chromatograph has four methods to separate different molecules: pressure, length of column, type of column, and temperature. By making use of all of these variables together, most compounds will separate well.

Pressure regulation is controlled at the injection port and can be varied to meet certain applications by a feature called EPC (Electronic Pneumatic Control). There are three modes EPC can be used to separate types of molecules in a sample: split, splitless, and pulsed split.

Split Mode: During a split injection, a liquid sample is introduced into a heated inlet where it vaporizes rapidly. A small amount of the vapor enters the column while the major portion exits from the split / purge vent. Split injections are primarily used for high concentration samples when you can afford to lose most of the sample out of the split / purge vent. It is also used for samples that cannot be diluted.

*Splitless mode:* In this mode, the purge valve is closed during the injection and remains so while the sample is vaporized in the liner and transferred to the column. At a specified time after the injection, the purge valve opens to sweep any vapors remaining in the liner out the split vent. This avoids solvent tailing due to the large inlet volume and small column flow rate. Since the entire sample gets transferred onto the column, this mode is primarily used for samples of low concentration.

Pulsed Split: The pressure pulse modes increase inlet pressure just before the beginning of a run and returns it to the normal value after a specified amount of time. The pressure pulse sweeps the sample out of the inlet and into the column faster, reducing the chance for sample decomposition in the inlet. This is helpful for large molecular weight compounds that tend to linger around in the inlet and thus tend to get purged in other EPC modes. This method can also help to increase sensitivity by placing a larger amount of sample on the column while decreasing the possibility of samples staying in the injection port and causing contamination.

The resolving power of a column can be dependent on the length of the column. The longer the column, the greater the resolving power. Longer columns allow for more interaction from each molecule in the sample with the stationary phase. Resolution between similar compounds with small differences can be achieved by increasing the length. Columns can be purchased in many lengths, but the growing trend is toward smaller columns since larger columns are more expensive and greater resolution by other factors can compensate for less resolution from the column.



Revision #19, Effective Date 10-20-2025

Capillary columns are a glass tube with a silianized coating containing different functional groups. Hundreds of different types of columns exist, each with a different type of functional group that will separate different compounds of interest. Some stationary phases can select for nonpolar, polar, and even compounds with lone pairs of electrons. The silianized layer thickness has a direct effect on the retention and elution temperature for each sample compound. Thicker films retain compounds longer by maximizing the amount of time the compounds spend in the stationary phase. Thinner films allow compounds to pass through the column faster, most likely with less separating ability.

Temperature variations in the injection port and on the column are important for separation by capillary gas chromatography. The injection port is usually set at 290°C, which vaporizes samples upon introduction to the GC. The injection port is not part of the column; it's a chamber where the liquid sample can change into the gas phase before entering the column. The oven, that contains the column, is relatively cool at a starting temperature of 190°C less than that of the injection port. This is hot enough to allow the volatile solvent to remain as a gas, but cold enough to cause the less volatile compounds to return to a liquid state. Once the sample becomes a liquid, it deposits itself on the column and won't migrate until the oven temperature heats up to the compounds boiling temperature. When the sample is once again in its gas phase, it travels through the column's stationary phase and mobile phase, jumping between the two. Separation has occurred through the difference in boiling points and through the amount of time different molecules spend in the stationary phase while traveling through the column.

By utilizing these four variables, separation of compounds can be achieved in most cases. Observation of the retention time is valuable in determining the identity of a compound, by comparison with a known standard. Changing just one of these variables can influence the retention time of a compound. The chromatograph should not contain wide peaks, since two or more compounds could resemble one peak. Ideally, sample concentration should be enough to give a single, narrow peak.

#### **References:**

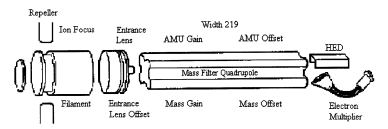
- Missouri State Highway Patrol Forensic Laboratory Chemistry Training Manual.
- 2. Hewlett-Packard GC/MS Product Software, August 1996.



Revision #19, Effective Date 10-20-2025

### **Appendix II - Basic Operation of the Mass Spectrometer**

In order to interpret a tune report, basic understanding of the MS is necessary. The following is a brief explanation of how the Electron Impact (EI) ionization method works. The molecules come off of the GC column and are subjected to electron bombardment, which causes them to fragment and become charged. A repeller is used to direct the ions to the focusing lenses in the ion source and then to the quadrupole mass filter. The mass filter allows selected ion masses to reach the detector. It separates ions based on their masses allowing only ions of a specific mass to reach the detector at a given time. The quadrupole filters by applying to each pair of quadrupole rods, a combination of radio frequency (Rf) and direct current (DC) voltages. One rod pair receives Rf voltage 180 degrees out-of-phase with the other pair, while an equal but opposite DC potential is applied to each rod pair. Under these conditions, at any particular set



of Rf and DC voltage values, only ions of a specific mass to charge (m/z) will traverse the length of the open space between the rods. All other ions are neutralized as they strike the surface of the rods. During a typical run, the MS scans

for masses ranging from 40 - 550 atomic mass units (amu). It scans for each mass unit in that range starting at the highest amu, working downward, throughout the duration of the run, with the exception of the solvent delay in which the MS is turned off. For instance, at the beginning of the scan, the mass filter selects only for masses of 550 amu, then it selects for masses of 549, and so on. This whole selection process takes place about three times a second. After each ion passes through the quadrupole, it is amplified by the electron multiplier, before reaching the detector. The detector counts the ions of each mass and plots the data on the mass spectrum (abundance vs. mass size). The quadrupole mass filter can select ions in two modes: Scan and SIM. Scan mode selects ions in the whole mass range specified, whereas SIM selects for specific mass units. The Scan mode has a lower sensitivity since most of the ions in a sample collide with the quadrupole rods. However, since samples are generally unknown, the filter mode utilized at the OSBI is the Scan mode to detect the entire spectrum of ions.

#### **Tuning**

Tuning is the process for optimizing the performance of the Mass Selective Detector (MSD). The goal of tuning is to maximize sensitivity while maintaining acceptable resolution (the ability to distinguish between a mass and its isotope), ensuring accurate mass assignment, and providing the desired relative abundances across the spectrum. The Mass Spec (MS) uses Perfluorotributylamine (PFTBA) because its mass spectrum has ions in the low (69), medium (219), and high (502) mass range. The mass spectrum of PFTBA is shown on the bottom of the tune report. The instrument looks specifically for masses 69, 219, and 502 in the spectrum of PFTBA, and plots these values along a mass axis (the x-axis of the mass spectrum). The instrument



Revision #19, Effective Date 10-20-2025

aligns its internal mass axis to match the PFTBA mass axis. Resolution is determined by the ability of the instrument to distinguish two peaks, one mass unit apart. This resolution is displayed on the tune report by the peaks in the upper left-hand corner. The peak graph displays the mass of the peak, the abundance of that ion, and the peak width at 50% of the height (Pw50), as shown in the upper left portion of the graph. The x-axis of this graph is the mass assignment. By increasing the Pw50, the area at the base of the graph increases, and a larger range is allowed for the selected mass assignment (i.e., 69, 219, or 502). If made too large, the base area could encompass a range more than its peak range and label the isotope mass with the selected mass. This would achieve greater sensitivity, but the resolution would be very low since it could not distinguish the selected mass from its isotope. It is for that reason that the peak width must be between 0.4 and 0.6. While viewing the tune report, look for clear separation between the selected mass and its isotope.

On the upper right of the printout, there are numerous parameters displayed. These parameters are automatically assigned while using autotune to optimize the MSD performance. These values can be manually changed by using manual tune, but is not recommended for normal use.

**Ion Pol:** This is the polarity of the field lens. A

positive field pushes the ions out of

the ion source. (5975)

Emission: The amount of current running

through the filament. The higher the current the greater the electron bombardment but decreases a filament life. Too low of a current will result in less ionization and reduced

sensitivity.

**ElEnrgy:** The electron energy of the electron

leaving the filament. (5975)

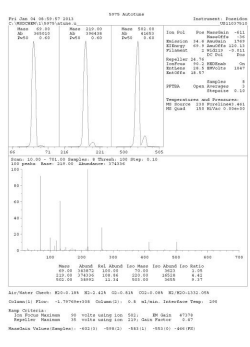
**Filament:** The MS contains two filaments in case one burns out.

**Repeller:** Sets the voltage of the repeller (part of the ion source). The repeller is a positive

Page **59** of **72** 

potential that repels the ions, pushing them out of the source. If the repeller is set too low, too few ions will leave the source, resulting in poor sensitivity and poor high mass response. If it is set to high, too many ions at too high a velocity will leave the source. This results in poor mass filtering and poor low mass

resolution.





Revision #19, Effective Date 10-20-2025

**IonFcus:** Sets the voltage of the ion focus lens (part of the ion source). Ion focus affects Ion

abundance. Generally, the offset is ramped during the tuning to find the ion focus

offset that results in the best ion abundance.

**EntLens:** Refers to the entrance lens gain, a value used to determine a mass dependent

voltage that is applied to the entrance lens. The entrance lens is the final lens through which ions pass before they enter the mass filter quadrupole. Typically, during tuning, the entrance lens voltage is ramped to find the setting that provides

the maximum abundance.

**EntOffs:** This is a constant voltage that is applied to the entrance lens. Increase the offset

to increase abundance of ions at low masses without substantially decreasing the

abundance of ions at high masses.

**PFTBA:** The status of the valve containing the PFTBA. This valve will open and close

automatically for tuning.

MassGain: Sets the value of the mass axis gain, which is a multiplicative factor used in the

equation to calibrate the mass axis. Mass gain adjusts the reported value of a given mass to the correct number. The mass that appears in a report has had a linear correction applied to it. This may be thought of as a calibration curve where the uncorrected mass is plotted along the x-axis and the reported mass is plotted along the y-axis. The calibration curve is a straight line with a slope that is proportional to the mass gain. Mass gain has a greater effect on mass assignments

at the high end of the mass scale than at the low end.

**MassOffs:** This is an additive factor used in the equation to calibrate the mass axis.

**AMUGain:** Atomic mass unit gain affects the width of the mass peak by adjusting the ratio of

DC voltage to RF voltage on the mass filter. A higher value gives narrower peaks,

but affects peaks at high masses more than those at low masses.

**AMUOffs:** This affects the width of the mass peaks by adjusting the ratio of DC voltage of the

mass filter quadrupole. A higher value gives narrower peaks at all masses.

Wid219: Affects the width of the mass peak at 219 amu. The value entered for this

parameter is approximately the value of the correction applied at mass 219. For instance, if a peak width adjustment has been performed and the values are: Mass 69 Pw0.60, Mass 219 Pw0.63, Mass 502 Pw0.60, then entering a value of -0.03 for the Wid219 parameter, followed by a peak width adjustment, should result in the

peak widths of all masses being set very close to 0.60 amu.



Revision #19, Effective Date 10-20-2025

**DC Pol:** Sets the polarity of the direct current applied to the quadrupole mass filter. This

parameter is set at the optimum polarity at the factory and should not be changed

for normal use.

**HEDEnab:** The High Energy Dynode sets the voltage to focus the ions into the detector, which

is located off-axis, hidden from photons and electrons coming from the source. The optimal HED voltage depends on the electron multiplier setting. Thus, the electron multiplier voltage is usually set first. Then the HED voltage is ramped to determine the setting that provides the greatest abundance. The older instruments assigned a value to this parameter, which use to be called X-ray lens, however the HP 5975 MSD does not have an X-ray lens and just indicates "on" or

"off". (5975)

**EMVolts:** The electron multiplier increases the abundance of all ions in the scan range going

to the detector.

Samples: The log2 of the number of samples to be taken and averaged at each mass during

a scan.

**Averages:** The number of profile scans to be averaged for each scan reported.

**Stepsize:** The mass axis increment used for a profile Scan. The larger the number, the faster

scans are taken, at a cost of resolution.

**Temperatures and Pressures:** 

MS Source/

**MSQuad:** Displays temperature settings for the Source and Quadrupole. (5975)

Foreline: The pressure between the rough pump and the diffusion pump. This area will

either state the pressure of the foreline if the MSD uses a diffusion pump or the

speed of the turbo pump. (5975)

**HiVac:** Displays the high vacuum pressure. (5975)



Revision #19, Effective Date 10-20-2025

On the Display of the mass spectrum of PFTBA, other parameters are listed:

**Scan:** 10.00-700.00 amu is the scan range during the tune. Typically, when drug samples

are scanned, this parameter is approximately 40-550 amu.

**Samples:** The log2 of the number of samples to be taken and averaged at each mass during

a scan. If the number is 8, log2 would be 256 scans.

**Threshold:** Abundance's below this value will be ignored for scanning. This determines what

signal will be accepted as peaks.

**Base:** Shows the base peak in the sample.

**Abundance:** Abundance of the base peak.

### **Tune Evaluation (Tune Eval)**

Tune evaluation is a way of verifying the performance of the MSD. First, it will evaluate the most current autotune (ATUNE.U) parameters and when the evaluation is complete, a system verification report is printed.

If all parameters of the autotune are within the predetermined limits, set by Agilent, they will be listed as "OK." If all parameters pass, the instrument can be used for casework. If any of the parameters fail, the reason for failure must be determined and corrected. The Autotune and Tune Evaluation must be run again and pass all parameters before casework can be analyzed on the instrument.

	System Verificati	on -	Tune	(Detector	Optimization)	Portion
Instrument N		:	Posei	don		
DC Polarity			Posit	ive		
Filament		:	2			
	uld be 69 or 219					Ok
Position of					69.00	Ok
Position of					219.00	Ok
Position of					502.00 70.00	Ok
	isotope mass 70				70.00	Ok
	isotope mass 220				220.00 503.00	Ok
Position of	isotope mass 503				503.00	
Ratio of mas	1sotope mass 503 s 70 to mass 69(0 s 220 to mass 219	.5 -	1.6%)		1.09	Ok
Ratio of mas	s 220 to mass 219 s 503 to mass 502 to 69 should be	(3.2	- 5.4	(8)	4.38 10.03 108.66 11.43	Ok
Ratio of mas	s 503 to mass 502	(7.9	- 12.	3%)	10.03	Ok
Ratio of 219	to 69 should be	> 40	and	18	108.66	Ok
Ratio of 502	to 69 should be	> 2.	4% and	i is	11.43	Ok
	ursor (<= 3%)				0.17	Ok
	cursor (<= 6%)				0.78	Ok
Mass 502 Pre	cursor (<= 12%)				1.35	Ok
	for a leak in the	sys	tem			
Ratio of 18					0.16	Ok
Ratio of 28	to 69 (<10%)				2.39	Ok
Electron Mul	tiplier Voltage				1047	Ok
Tune por	tion of System Ve	rifi	cation	passed.		

#### **References:**

- 1. Missouri State Highway Patrol Forensic Laboratory Chemistry Section Training Manual.
- 2. Hewlett-Packard GC/MS Product Software, August 1996.
- 3. www.agilent.com



Revision #19, Effective Date 10-20-2025

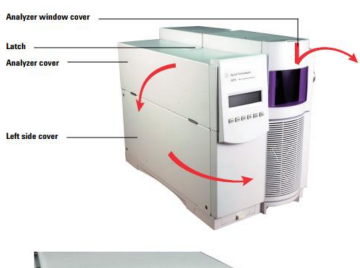
## Appendix III - Helpful Tips for The Mass Spectrometer

#### Pump down/Filament/Vent

 In the Instrument Control view, select Instrument, select MS Vacuum Control to display the Vacuum Control dialog box or select View, select Tune Vacuum Control, select Vacuum select MS Vacuum control

#### 2. Click Vent

- 2.1. Follow the instructions presented on the screen, and wait until the vent cycle is completed.
- 2.2. Let the rough pump cool off approximately 45 minutes if needed unplug the pump
- 2.3. Close the software and turn off the MS portion on the instrument
- 3. Turn the vent valve counterclockwise only 3/4 turns or until you hear the hissing sound of air flowing into the analyzer chamber





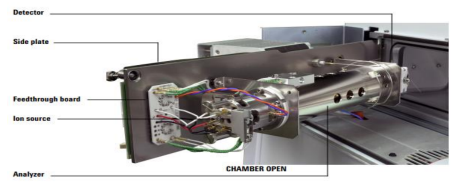
Page **63** of **72** 



Revision #19, Effective Date 10-20-2025

4. Disconnect the side board control cable and the source power cable from the side board. Loosen the side plate thumbscrews if they are fastened and gently swing the side plate out





- 5. Remove the source from the analyzer
- 6. Disassemble the source and collect the parts to be cleaned. Clean using microgrit /sandpaper. Sonicate metal pieces using methanol approximately 15 minutes. Ensure microgrit is cleaned off before reassembling the source.

**Side note:** Ensure the new filaments are placed the correct way!



- 7. Reinstall the source into the analyzer and close the side plate
- 8. Reconnect the side board control cable and source power cable to the side board
- 9. Plug the rough pump back in



Revision #19, Effective Date 10-20-2025

- 10. Make sure the vent valve is closed turn knob clockwise
- 11. Press the Power on button on the front of the MS while pressing on the side plate to ensure a good seal
- 12. Start the Masshunter Data Analysis program. In the Instrument Control view, select Instrument, select MS Vacuum Control to display the Vacuum Control dialog box or select View, select Tune Vacuum Control, select Vacuum select MS Vacuum control
- 13. Select Pump Down
- 14. Allow ample time for instrument to pump down

#### **Switch Filaments**

- 1. From the Instrument Control menu, select View
- 2. Select Parameters next select Manual Tune
- 3. Enter the filament number
- 4. Click Done and save the atune file or select Save Tune Parameters from the File menu

#### Manual Tune for air leak

- 1. From the Instrument Control menu, select View
- 2. Select Parameters next select Manual Tune
- 3. Select the **Scan** tab of the **Manual Tune** dialog box and set the Scan Mass Range to Low m/z 10 to High m/z 100 also check that the **PFTBA** is set to **Closed** in the **Parameters** section.
- 4. Click Scan, look at the window labeled scan
- 5. Do not save atune

#### **Search Drug library**

- 1. From the Instrument Control menu select View, select MSD Chemstation Data Analysis
- 2. Select View next select Parametric Retrieval
- 3. Enter in the library you would like to search default: c:\Database\CrDM.I Note: Other libraries are available to be searched
- 4. Choose your search option in the side panel

5975 Series MSD Operation Manual for MassHunter (agilent.com)



# **Appendix IV - Mass Peaks of Common Contaminants**

Mass(es)	Compound General Classification	Potential Source
18, 28, 32, 40, 44	Air	H <sub>2</sub> O, N <sub>2</sub> , O <sub>2</sub> , Ar, CO <sub>2</sub>
18	Cleaning Solvents	Water
31		Methanol
47, 83, 85		Chloroform
77		Benzene or Xylenes
91,92		Toluene
105,106		Xylenes
43, 58		Acetone
85		Freons
73, 147, 207, 222, 281,	Dimethylpolysiloxane	Septum or Column
295, 341, 355, 429	2 meeny peryonoxane	bleed
41, 43, 55, 57,	Hydrocarbons	Fingerprints or
71, 85, 99	Tiyurocarbons	pump oil
149	Phthalates	Plasticizers in tubing,
		vials, caps, samples

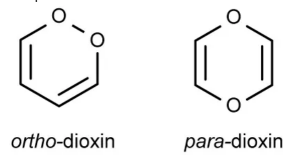


Revision #19, Effective Date 10-20-2025

### Appendix V – Isomers

**Structural isomers:** a form of isomers in which molecules with the same Molecular Formula have different bonding patterns and atomic organization.

- Structural isomers share the same chemical formulas, but their atoms are arranged differently.
- Examples include ortho- para- meta-



**Stereoisomers:** a form of isomers in which molecules with the same Molecular Formula have molecular bonds which are always in the same order and only differ in spatial arrangement.

- Isomers that have the same number of the same kinds of atoms but differ in the orientation of their atoms in space. These isomers differ in chemical and physical properties.
- There are two types of stereoisomers: enantiomers and diastereomers.

**Enantiomers or Optical isomers:** 2 isomer molecules that are chiral, and are <u>mirror images</u> of each other; a pair of chemical compounds whose molecular structures have a <u>nonsuperimposable</u> mirror-image relationship to each other



Revision #19, Effective Date 10-20-2025

**Diastereomers or Geometric isomers:** 2 stereoisomers that are not mirror images; a stereoisomer of a compound having two or more chiral centers that is not a mirror image of another stereoisomer of the same compound

diastereomers



## **Mock Trial Evaluation Form**

Analyst		Score	e	
Reviewer		Date	e	
Please rate the trainee's performance during the Mock	Trial: Excellent (3)	Good (2)	Fair (1)	Poor (0)
Courtroom demeanor and appearance				
Ability to convey information in an understandable manner				
Poise and professionalism during direct examination				
Poise and professionalism during cross examination				
Use of court exhibits/visual aids (if applicable)				
Testimony based upon scientific principles				
Exhibition of knowledge of OSBI testing procedures				
Explanation of results				
Remarks/Comments/Suggestions/Explanation for Poor	Ratings:			



Revision #19, Effective Date 10-20-2025

## **Additional Reading/Training**

This section is intended to list articles/books/journals/etc. that are required reading, additionally training courses can be listed as well.

Date	Literature/Training



## **Approval**

Technical Manager  Michella Carter	Date	9/23/25
Criminalistics		
Division		
Director Xumu Xu	Date	09/23/2025
000		
Comments		



## History

Revision	Issue Date	History
19	10-20-2025	Drug Quality System Overview: Updated the Literature
		section from Review OSBI CSD QM 6.1 to Read Physical
		Evidence Quality Procedures Manual 2.1 Evidence Handling.
		Weights and Measures: Added to Discussion for
		Uncertainty of Measurement "Where should uncertainty be
		added to the report for marijuana cases that include both grams and pounds."
		Thermometers: Updated the Literature section from
		Review OSBI CSD QM 6.4 to Read Physical
		Evidence Quality Procedures Manual 2.4, Evidence
		Refrigerator and Freezer Maintenance.