1. **Scope**

1.1. This document establishes the procedures for quantifying heroin in samples containing heroin for use in case work and reporting quantifiable results of heroin in test samples using Gas Chromatography Flame Ionization Detector (GC-FID) instrumentation.

2. **Background**

2.1. This method is based off the document provided by the Drug Enforcement Agency (DEA) method for *Quantitation of Heroin* (2016) using GC-FID. This document provides guidance and is in support of the Forensic Chemistry Unit Quality Assurance Manual (QAM) and conforms to the requirements of the accreditation standards under ISO/IEC 17025 (current revision) and the supplemental standards.

3. **Safety**

3.1. Reagent Toxicity: Personnel should refer to the appropriate SDS for solvents and reagents used during analysis for any specific safety requirements.

3.1.1. For a complete review of required Health and Safety regulations of the PHL, see *DOM13 DFS Health and Safety Manual*.

3.2. Protective Equipment: Personnel should wear personal protective equipment (PPE) including: lab coat, gloves, and safety goggles when carrying out standard operating procedures.

3.2.1. Wear vinyl or nitrile gloves when handling these chemicals to prevent
absorption through the skin. If any chemicals are spilled onto gloves, discard gloves into hazardous waste.

3.3. Training: Formal training in use of instruments and software is necessary.

3.4. Personal Hygiene: Universal Precautions shall be followed. Care should be taken when handling chemicals or any biological specimen. Routine use of gloves and proper hand washing should be practiced.

3.4.1. Refer to DOM13 – DFS Health and Safety Manual.

3.5. Disposal of Waste: Waste materials shall be disposed of in compliance with laboratory, Federal, state, and local regulations. Solvents and reagents should always be disposed of in an appropriate container clearly marked for waste products and temporarily stored in a chemical fume hood.

3.5.1. Consult DFS Safety Officer for proper procedures.

4. Materials / Equipment Required

4.1. Reagent Grade Acetonitrile (ACN), Methanol (MeOH), and Chloroform (CHCl₃), or higher purity.

4.2. Ultra-High Purity Helium tanks

4.3. Hydrogen generator or tanks for the FID

4.4. Gas Chromatography Flame Ionization Detector (GC-FID), fully assembled, including: solvent vials, injection syringe, and other consumables.

4.5. Glassware:

4.5.1. 250 mL volumetric flask (Class A)
4.5.2. 100 mL volumetric flask (Class A)
4.5.3. 1-10 mL volumetric pipettes (Class A)
4.5.4. Other volumetric glassware (Class A), as appropriate

4.6. Binder/Folder for Standard results, or electronic equivalent

4.7. Maintenance Logbook, or electronic equivalent

5. Standards and Controls

5.1. Standards may come from Cayman Chemical or the Drug Enforcement Agency (DEA), or other, approved vendor or provider.

5.2. Heroin hydrochloride (HCl) (solid powder)
5.3. Tetracosane (solid powder)

5.4. Mannitol (solid powder)

5.5. Internal Standard Solution (ISTD), 0.40 mg/mL tetracosane (exact value needed):

5.5.1. About 100mg tetracosane in a 250 mL volumetric flask, dilute to mark with chloroform/methanol (9:1). Other volumes are acceptable as long as ratio is maintained.

5.6. Heroin HCl Stock Solution, about 2mg/mL (exact value needed using purity):

5.6.1. About 200mg Heroin HCl in tared, 100mL volumetric flask. Other volumes are acceptable as long as ratio is maintained.

5.6.2. Dilute to mark with ISTD solution.

5.7. Heroin HCl Calibration Solutions: (dilute to mark with ISTD solution; exact value needed using purity):

5.7.1. Cal 1: 1:10 about 0.2 mg/mL
5.7.2. Cal 2: 2:10 about 0.4 mg/mL
5.7.3. Cal 3: 3:10 about 0.6 mg/mL
5.7.4. Cal 4: 4:10 about 0.8 mg/mL
5.7.5. Cal 5: 5:10 about 1.0 mg/mL
5.7.6. Cal 6: 6:10 about 1.2 mg/mL
5.7.7. Cal 7: 7:10 about 1.4 mg/mL
5.7.8. Cal 8: 8:10 about 1.6 mg/mL
5.7.9. Cal 9: Heroin HCl Stock Solution (about 2.0 mg/mL)

6. Calibration

6.1. Response to Heroin HCl Calibration Solution

6.1.1. An average of five responses are taken at each calibration level

6.1.2. The average response of each concentration level is compared to the actual concentration, with an average sensitivity calculated at each level (sensitivity is the response per concentration).

6.1.3. For the concentration curve to be acceptable, the average sensitivity of each calibration level shall be within 5% of the overall average sensitivity.

6.1.4. While in use for casework, approved calibration Excel worksheets will be saved electronically.
6.1.5. Once a curve is no longer suitable for casework, the respective Excel worksheet will be archived in the appropriate electronic folder.

6.1.6. This calibration curve will be run at least once per year or as needed (based on column environment or method changes) to evaluate the suitability of the column for quantitation and a re-establishment of working range.

6.2. Monthly/Weekly Calibrant Run with Samples

6.2.1. A calibrant of heroin standard at approximately 1 mg/mL and two quality control standards (low and high) will be run for each sequence at least once each month that heroin purity analysis is performed.

6.2.2. The calibrant will be used to establish the calibration curve for that month’s analysis (or until another calibrant is run).

6.2.2.1. The origin and the response from the calibrant will be the two points used to determine the new calibration curve between calibrant runs.

6.2.3. The calibrant response will be monitored in an electronic control chart.

7. Procedures

7.1. Method Parameters: Gas Chromatograph-Flame Ionization Detection (GC-FID)

7.1.1. Split mode (60:1, split flow 60 mL/minute)

7.1.2. Column: HP-5, 12mx0.20mm inner diameter (I.D.) x 0.33 µm film thickness, 5% phenyl methylpolysiloxane stationary phase, or equivalent.

7.1.3. Column Flow Program: 1.0 mL/minute for 2.5 minutes, ramp 45 mL/minute to 4.5 mL/minute, hold for 1.0 minute.

7.1.4. Oven Program: 270ºC for 2.5 minutes, ramp 45ºC/minute to 295ºC, hold 1.0 minute.

7.1.5. Inlet (injector) Temperature: 280ºC (pressure 12.665 psi, total flow 64 mL/minute, septum purge flow 3 mL/minute)

7.1.6. Detector: 280ºC at 50Hz

7.1.7. Carrier Gas: Helium

7.1.8. Run time: 4.05 minutes

7.1.9. Injection volume: 1 µL

7.1.10. Injection Solvent: Chloroform / Methanol (9:1)

7.2. Acceptable Method Parameter Variations (make mention in case notes):
7.2.1. Inlet temperature may be changed from about 270ºC to 290ºC for better improved chromatography, as per analyst’s discretion.

7.2.2. Increase flow and temperature ramp at 2.5 minutes to remove late eluting compounds (e.g., diltiazem, noscapine, ...), as per analyst’s discretion.

7.2.3. Sample preparation solvent of chloroform or chloroform/methanol (9:1), as per analyst’s discretion.

7.2.4. Data sampling rate may be changed from 20 to 50Hz, as per analyst’s discretion.

7.2.5. Each linearity concentration may be injected in lowest-to-highest or highest-to-lowest order, as per analyst’s discretion.

7.2.6. Accuracy and recovery may be calculated from either a 3-point or 9-point curve, as per analyst’s discretion.

7.3. Quality Control Standards – Two Quality Control Standards are made and evaluated over time to assess performance of the method. The bulk QC Heroin mixture is made:

7.3.1. About 50mg Heroin HCl (>99% purity) added to about 850mg sucrose or mannitol.

7.3.2. Mix contents and grind in mortar

7.3.3. (Optional) Filter through approximately 20 mesh.

7.3.4. Quality Control (QC) standards (for example):

7.3.4.1. QCLow – About 75mg bulk QC Heroin Mixture, added to 10mL volumetric flask, dilute to mark with ISTD solution

7.3.4.2. QCHigh – About 300mg bulk QC Heroin Mixture, added to 10mL volumetric flask, dilute to mark with ISTD solution

7.3.4.3. Filter prior to injection.

7.4. Unknown Samples – Accurately weigh the sample and dissolve in the ISTD solution. Document sample weight in case notes.

7.4.1. Add sufficient quantity to result in a concentration that is within the working range of this method. Example: weigh approximately 10mg of sample and dilute with 2mL of ISTD.

7.4.2. Filter prior to injection.

7.5. **Acceptance Parameters.** – The following are the acceptance parameters for the GC-FID.

7.5.1. GC-FID Peak Quality
### GC-FID PARAMETERS

<table>
<thead>
<tr>
<th>Acceptance Criteria</th>
<th>Detail</th>
</tr>
</thead>
<tbody>
<tr>
<td>Retention Time</td>
<td>Retention time of analyte peak must match within 2% of standard.</td>
</tr>
<tr>
<td>S/N* Cut Off</td>
<td>Analyte TIC** must be more than three (3) times greater than noise.</td>
</tr>
<tr>
<td></td>
<td>(S/N=signal-to-noise ratio)</td>
</tr>
<tr>
<td>Peak width resolution</td>
<td>Analyte peak must be base peak resolved, as evaluated by the analyst.</td>
</tr>
</tbody>
</table>

*S/N = Signal-to-Noise Ratio; **TIC = Total Ion Chromatogram

### 7.5.2. Calibration Curve Acceptability

<table>
<thead>
<tr>
<th>Acceptance Criteria</th>
<th>Detail</th>
</tr>
</thead>
<tbody>
<tr>
<td>Check standard falls outside of tolerance</td>
<td>Tolerance set to 5%</td>
</tr>
<tr>
<td>Calibration curve still valid</td>
<td>Calibration curve (two-point recalibration) is applicable for all casework within one month of creation</td>
</tr>
<tr>
<td>Two samples of the same item are within 10%</td>
<td>Absolute difference in quant of two samples of an item must be under 10% in order to be acceptable</td>
</tr>
<tr>
<td>Range of &gt;10% between samples requires third quantitation</td>
<td>Analyst may average the sample concentrations or report the lowest value</td>
</tr>
</tbody>
</table>

### 7.5.3. QC Check Acceptance
7.5.3.1. Positive control checks shall be within 10% of each other to be acceptable.

7.5.3.2. If a QC check value does not fall within this range, a new standard solution may be prepared and injected.

7.5.3.3. If more than one QC check value does not fall within this range, a new calibration curve shall be prepared and used.

7.6. Sample Runs

7.6.1. Prepare two separate samples (three if necessary) for casework. Record weight values using a five place balance (reading down to 0.00001g). Each sequence shall include a negative control, calibrant, and two QCs (low and high).

7.6.2. Filter and inject the solution into the GC-FID.

7.6.3. From the generated data, divide the heroin peak corrected area by the internal standard’s peak area to calculate the area ration.

7.6.4. Using the two-point recalibrated calibration curve (from section 6.2), solve for the heroin concentration.

7.6.5. To calculate the percent heroin:

7.6.5.1. Slope of Calibration Curve = (Area of calibrant peak) / (Area of internal standard calibrant peak x Concentration of Calibrant (mg/mL))

7.6.5.2. Percent Purity = 100% x (Area of sample heroin peak) / (Slope of Calibration Curve x Area of sample internal standard peak x Total Concentration of Sample (mg/mL))

7.7. Control Chart Maintenance

7.7.1. As appropriate, the significant parameters appropriate for the identification of individual substances shall be recorded in the laboratory control chart for GC-FID. Critical pieces of information include peak retention time and QC purity values.

8. Sampling

8.1. Perform sampling plan as covered under section 8 of FCS02 – SOP for General Laboratory Procedures for FCU. Note: Heroin quantitation will not be performed on samples that fit any of the following criteria:

8.1.1. The total net weight of the sample is less than 100 mg before analysis (or, in the case of multiple homogenous units, estimated net weight of all possible units).

8.1.1.1. A composite of multiple samples may be necessary to meet the minimum weight required to perform quantitative analysis
(100 mg). In cases where a composite shall be made, the chemist will first test each of the selected samples with a screening technique prior to making a composite (Category A, B, C). The number of items composited shall be documented in the case notes.

8.1.2. The heroin is a very minor component in the sample, as evaluated by the analyst (based on the relative abundance in GCMS. The estimated cutoff is less than 5%).

8.2. After quantitation, the heroin result will not be reported if the results fit any of the following criteria:

8.2.1. The purity determination is below the detectable limit.

8.2.2. The purity determination is below the uncertainty value (the reported uncertainty value).

8.3. When heroin is detected but not quantitated, a note will be made on the report stating the reason why quantitation was not performed or why a quantitation result was not reported.

8.4. Deviations are acceptable but shall be recorded in case notes and follow agency policy of deviations.

9. Calculations

9.1. Linearity Evaluation of Calibration Standards

The following is an example calculation and evaluation of response values for the 9 calibrators used in this method, using Excel.
9.2. Regarding the quantitation, or purity, analysis of test items for heroin, the standard result will be assuming a salt form of “heroin hydrochloride” or “heroin HCl.” The calibration curve on the gas chromatogram flame ionization detector (GC-FID) instrument will generate percentages assuming the heroin HCl. In instances where a purity is requested assuming a heroin base, or vice versa, a conversion factor may be used. This value is the ratio of the molecular weight of the heroin base to heroin hydrochloride (hydrated), or:

9.2.1. Calculate Heroin Base from Heroin Hydrochloride

\[
\text{9.2.1.1. Purity (heroin base)} = \text{Purity (heroin HCl)} \times \left(\frac{369.417 \text{g/mol heroin}}{423.8902 \text{ heroin HCl [H2O]}}\right)
\]

\[
\text{9.2.1.2. Purity (heroin base)} = \text{Purity (heroin HCl)} \times 0.871492
\]

9.2.2. Calculate Heroin Hydrochloride from Heroin Base

\[
\text{9.2.2.1. Purity (heroin HCl)} = \text{Purity (heroin base)} \times \left(\frac{423.8902 \text{ heroin HCl [H2O]}}{369.417 \text{g/mol heroin}}\right)
\]

\[
\text{9.2.2.2. Purity (heroin HCl)} = \text{Purity (heroin base)} \times 1.147457
\]
10. Uncertainty of Measurement

10.1. Uncertainty of measurement is assessed as per the validation of this method. See corresponding validation per instrument.

10.2. See FCS21 – Procedure for Uncertainty in Measurement, for purity and weight uncertainty calculations.

11. Limitations

11.1. Not applicable

12. Documentation

12.1. Case notes
12.2. Instrument Logbooks

13. References


13.3. FCS02 – SOP for General Laboratory Procedures for FCU.

13.4. FCS21 – Procedure for Uncertainty in Measurement


13.7. American National Standards Institute National Accreditation Board Supplemental Requirements (current revision)