

Centre of Forensic Sciences Investigators and Submitters

Technical Information Sheets Toxicology

November 2023

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Introduction

The Toxicology Section performs analyses on biological samples (e.g., blood, urine, liver) to determine the absence/presence/concentration(s) of drugs, including alcohol and poisons.

This document is intended as a convenient investigative reference but should not be relied upon as definitive or exhaustive. Please contact the Centre of Forensic Sciences (CFS) Toxicology Section for assistance with questions of an analytical or toxicological nature by e-mail or telephone (647-329-1400 or 647-329-1430). When calling please ask for the appropriate coordinator:

Coroner's Coordinator:

CFSToxicologyCoronerCoordinator@ontario.ca

Criminal Coordinator:

toxcrim@ontario.ca

Examination Strategy and Capability

The screening methods employed in the Toxicology Section are:

- Gas Chromatography/Mass Spectrometry (GC/MS)
- 2. Immunoassay (IA)
- 3. Head-Space GC analysis for volatiles
- 4. Quadrupole Time-of-Flight MS (QTOF)

The targeted/quantitation methods employed in the Toxicology Section are:

- GC
- 2. Liquid Chromatography (LC), LC-MS/MS
- 3. Head-Space GC analysis for volatiles

Capabilities of screening methods are presented in Appendix 1. While these screening methods have wide-ranging capabilities, not all drugs may be reliably detected. Appendix 2 contains a list of compounds that may not be identified by the screening methods but may be detected/quantitated by targeted methods. Many of the compounds contained in this list will not be tested for unless specifically requested. If use of a specific drug is known or suspected and is relevant it should be noted in the case synopsis.

The examination strategy, i.e., determining which tests will be performed in a case, is informed by a variety of sources including case type, case history, nature of submitted samples, analytical protocols and capabilities, and discussions with clients. The initial toxicological analyses conducted for a variety of case types are presented in Appendix 3.

Urgent Cases

Requests for expedited analyses must meet specific criteria before being accepted as an urgent case. This process requires authorization by Toxicology Section management.

Examination

All items are visually examined on receipt to check the seal numbers (if present), the contents, and the integrity of the packaging.

Instrumentation

Chromatography: Gas Chromatography (GC); Liquid Chromatography (LC)

Chromatography is an analytical technique used to separate compounds based on their chemical and structural properties. GC uses a pressurized gas, while LC uses a pressurized liquid, in the separation of compounds.

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Immunoassay (IA)

IA detects compounds in biological fluids using a reaction of an antibody or antibodies to its antigen (i.e., the drug). This technique is primarily a screening technique; however, some IA methods are semi-quantitative, e.g., acetaminophen.

Mass Spectrometry (MS)

MS detects, identifies, and quantitates compounds. An MS can be coupled with a GC or an LC.

Quadrupole Time-of-Flight-MS (QTOF)

QTOF detects and identifies compounds. A QTOF is coupled with an LC.

Tandem MS (MS/MS)

MS/MS detects, identifies, and quantitates compounds and is commonly coupled to a GC or LC.

Ultraviolet and Visible (UV/VIS) Spectrophotometry

UV/VIS spectrophotometry identifies and/or quantitates a drug based on its UV and/or visible light-absorbing properties.

Carbon Monoxide

Carbon monoxide is analyzed by visible spectrophotometry. Results are expressed as % carboxyhemoglobin saturation.

Interpretation

Quantitative results may be expressed as 1) a concentration or 2) as < or > a concentration, e.g., when sufficient for interpretation. Blood ethanol interpretations provided in reports are generally limited to cases in which the detected concentration may be associated with fatalities, may be influenced by post-mortem artefacts, may have toxic interactions with other drugs, or in the case of motor vehicle collision, associated with impairment.

Measurement Uncertainty

Measurements made with all scientific instruments are associated with variability. No measurement is exact but is an estimate of the true value. Calculation of measurement uncertainty (MU) employs statistical methods to determine the range of values within which the quantitative result is likely to reside. The MU provides a reasonable estimate of the variability associated with the analytical method and is based on the analysis of matrix-matched quality control samples. A minimum of 10 such analyses are used. The MU is calculated with a confidence of 95.45 per cent using a k-factor based on the degrees of freedom as determined by the Student's t-test and the standard deviation of the associated quality control data. The MU is expressed in the same units in which the quantitative result is reported, e.g., ng/mL, mg/L and is reported as: quantitative result \pm MU.

Limitations

The focus of this laboratory is drug toxicity. Clinical blood/urine chemistry analysis, e.g., electrolytes, cell counts, gas saturation, creatinine, is not performed. Analysis for antiepileptic drugs is limited to determining drug toxicity, when warranted, based on case history. This laboratory does not have validated methods to analyze some sample types, e.g., oral fluid, hair, bile, muscle, brain tissue. There are a variety of analytical issues that may prevent the detection of some of the drugs that this laboratory is commonly capable of detecting, which include:

- matrix effects
 - degree of putrefaction
 - type of sample (e.g., splenic blood)
 - post-mortem interval
 - storage conditions
- volume of sample submitted
- low concentration of the drug/sensitivity of the method

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Conversely, some novel, or rarely encountered, drugs not listed in Appendix 1 may be identified by the GC/MS or QTOF screens. In this case, analytical reference material would be acquired (if available) then analysed to confirm identity. There are drugs/compounds for which the CFS Toxicology Section does not have a method, examples of which are provided in Appendix 4.

Glossary

Abbreviations

Analytical results are reported in terms of mg/100 mL, mg/L, or ng/mL, as shown below:

g gram
mg milligram
ng nanogram
L litre
mL millilitre

Breakdown Product

A compound produced either inside or outside the body that may or may not be pharmacologically active.

Carboxyhemoglobin saturation

The percentage of hemoglobin bound by carbon monoxide.

Central Nervous System Depression (CNS depression)

A lowering of the functional activity of the brain and/or spinal cord. Depression of the respiratory and the cardio-regulatory centres are most relevant toxicologically.

Confirmation

The process of verifying the presence of a drug by replicate analysis using the same or different analytical technique(s). Confirmation of an immunoassay result is achieved using a more specific analytical technique.

Coroner's Case Analytical Summary

Contains analytical results with the fatal reference and limitations. The Coroner's Case Analytical Summary is accompanied by an Interpretive Guide with information specific to this report type.

Detected

The drug has been identified in the sample. Identification is based on criteria specific to the analytical technique.

Fatal Reference

A minimum drug concentration at which death has been reliably reported in the forensic literature.

Inconclusive

The presence or absence of a drug could not be determined.

Metabolite

The product of enzymatic conversion of a drug within the body to a different compound that may or may not be pharmacologically active.

No [other] significant findings by a [method name(s)]

This comment is inserted to provide a reference to the methods that were used. Appendices 1 and 5 can be used to identify compounds not listed and that were either not detected or the results were deemed to not be toxicologically significant, e.g., caffeine or nicotine. This may also apply to endogenous compounds, e.g., acetone < 2 mg/100 mL.

Not Detected

The drug is either not present or is present but at an amount that cannot be discerned from other constituents in the sample.

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Post-mortem redistribution

A phenomenon that refers to a change (either an increase or a decrease) in blood drug concentration after death; post-mortem redistribution may occur regardless of sampling site but is commonly observed as increased drug concentrations in heart blood as compared to femoral blood.

Putrefaction

The decomposition of organic material that involves micro-organisms.

Report

Contains a comprehensive summary of analytical results accompanied by interpretative conclusions.

Tentative

A drug has been identified by a non-specific screening method (e.g., immunoassay) but has not been confirmed. Further analysis would be required to positively identify the compound.

Therapeutic

The detected drug concentration is generally considered to not be toxicologically significant. The use of this term does not imply clinical efficacy.

Traces

The drug was detected at a concentration below that which can be reliably quantitated. The use of this term does not imply clinical efficacy.

Unconfirmed

A drug has been identified by a single procedure but not quantitated or confirmed by a second analysis. Unconfirmed findings may or may not be toxicologically significant.

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Appendix 1 - Screening Methods

Drugs that can be reliably detected by screening methods

GC/MS Screen

Α cotinine cyclobenzaprine² alpha-pyrrolidinovalerophenone (α-PVP) cyproheptadine1 acetyl fentanyl2 amantadine1 amitriptyline² desipramine² amlodipine² dextromethorphan² amoxapine² dextrorphan* amphetamine² diazepam² amphetamine (4-fluoro) diazepam (nor)2 anabasine dibucaine dihydrocodeine atomoxetine atropine/hyoscyamine diltiazem² diltiazem (desacetyl)2 dimethyltryptamine benzocaine diphenhydramine² benzofuran (6-(2-aminopropyl, 6-APB) doxepin² benztropine1 doxylamine² benzylpiperazine (BZP) Ε bromo-dragonfly brompheniramine² ephedrine*2 bupivacaine1 estazolam bupropion² etizolam² butylone/ethylone ethylone/butylone butyryl fentanyl x-fluoroamphetamine caffeine1 fluoxetine² carbamazepine² fluoxetine (nor)2 cathinone (cath) flurazepam² n-ethyl-cath flurazepam (n-desalkyl)² 4-fluorometh-cath fluvoxamine² 3-methoxymeth-cath Н 4-methyleth-cath meth-cath haloperidol1 hydrocodone² chlorcyclizine hydroxychloroquine chlordiazepoxide² chloroquine hydroxyzine¹ chlorpheniramine² chlorpromazine¹ cisapride ibogaine imipramine² citalopram*2 clomipramine² Κ clonidine1 ketamine² clozapine² cocaethylene cocaine² lamotrigine² codeine² laudanosine

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levamisole phenmetrazine lidocaine phentermine1 Ioratadine piperazine, 1-3 chlorophenyl (mCPP) loxapine² piperazine, trifluoromethylphenyl (TFMPP) p-fluorofentanyl M p-methoxyamphetamine (PMA)2 maprotiline1 p-methoxymeth-amphetamine (PMMA) meclizine1 procaine1 mefloquine1 prochlorperazine² meperidine² procyclidine1 meperidine (nor)2 propoxyphene² mephedrone² propranolol2 mepivacaine1 pseudoephedrine*2 methadone² Q methamphetamine² quetiapine² methamphetamine (4-fluoro) methedrone quinidine1 methotrimeprazine² R methylenedioxyamphetamine (MDA)² methylenedioxyethylamphetamine (MDEA)2 ropivacaine methylenedioxymethamphetamine (MDMA)² S 3,4-methylenedioxypyrovalerone (MDPV)² scopolamine (hyoscine)1 methylone² sertraline2 methylphenidate² metoclopramide1 strychnine¹ metoprolol² Т midazolam² tapentadol mirtazapine² terbinafine moclobemide1 ticlopidine tramadol² nicotine1 trazodone² trihexphenidyl2 nortriptyline² trimethoprim 0 trimipramine² triprolidine² olanzapine² orphenadrine² oxybutynin¹ oxycodone² valeryl fentanyl varenicline venlafaxine2 paroxetine² venlafaxine (O-desmethyl)² verapamil² pentadrone pentazocine² X pentoxyphylline² pentylone xylometazoline phenacetin Ζ phencyclidine (PCP)2 zolpidem²

zopiclone breakdown product

phenethylamines (2C-B, 2C-B-Fly, 2C-T-7, PEA)

Senior Manager, Toxicology

pheniramine²

Authorized:

QTOF Screen

The QTOF screen is a powerful and sensitive method that can reliably detect the drugs included in the following methods (details are listed in Appendices 5 and 6):

- LC-MS/MS Mix 2
- LC-MS/MS Mix 3 (except carfentanil)
- LC-MS/MS Mix 4
- LC-MS/MS Mix 5 (except: diflunisal, furosemide, ibuprofen, salicylate, vigabatrin)

In addition, the QTOF screen can identify psilocin. The list of drugs potentially identifiable by QTOF is too extensive to list within this document. For questions about a specific drug not listed, please contact the appropriate case coordinator.

*The GC/MS screen and QTOF screen are not capable of distinguishing racemates, therefore compounds such as dextrorphan/levorphanol, citalopram/escitalopram and ephedrine/pseudoephedrine cannot be separated. Similarly, the GC/MS screen cannot distinguish between 2-fluoroamphetamine, 3-fluoroamphetamine, and 4-fluoroamphetamine.

Immunoassay Tests (known cross-reactivity)

Barbiturates:

amobarbital² butalbital² pentobarbital² phenobarbital² secobarbital²

Head-space GC-FID analysis for volatiles (screen and quantitation)

acetone methanol

ethanol n-propanol (qualitative)

isopropanol

Volatile screen (qualitative only)

difluoroethane acetone propane dichloromethane butane methyl ethyl ketone 1,1,1,2-tetrafluoroethane isopropyl alcohol isobutane ethyl acetate toluene acetaldehyde diethyl ether methanol chloroform dimethyl ether ethanol gasoline

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Appendix 2 - Drugs Requiring Targeted Analysis

Compounds that may not be identified by screening methods but might be detected and/or quantitated by targeted methods.

A furosemide² S

antipyrine (phenazone)¹ I sufentanil¹

C ibuprofen² T

carbaryl¹ M toluene³

carbon monoxide⁴ methaqualone¹ **V**

mexiletine¹
valproic acid³

P
vigabatrin²

diflunisal² phenyltoloxamine¹ **F** physostigmine¹

fenfluramine¹ formic acid³

Methods used for the quantitation of compounds identified in the preceding appendices are denoted as follows:

¹GC-NPD ³GC-FID

² LC-MS/MS ⁴ Visible spectrophotometry

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Appendix 3 - Initial Analyses by Case Type^a

Alcohol-impaired driving: Ethanol

Attempted murder:dependent upon case historyConfirmation of ketoacidosis:Ethanol (includes acetone), BHB

Death of child < 5 years of age Ethanol, QTOF Screen, LC-MS/MS Mix 3, IA cannabinoids, IA acetaminophen

Drug-impaired driving: QTOF Screen, IA cannabinoids, UDM, GHB

Fatal motor vehicle collision (driver) and aviation death: Ethanol, QTOF Screen, LC-MS/MS Mix 3, IA

cannabinoids, COb

Fire-related death^c: CO (whole blood required)

Homicide: Ethanol, QTOF Screen, LC-MS/MS Mix 3, IA cannabinoids **Mandatory inquest:** Ethanol, QTOF Screen, LC-MS/MS Mix 3, IA cannabinoids

Possible drug-related death: Ethanol, QTOF Screen, LC-MS/MS Mix 3

Rule Out/exclusionary Toxicology: Ethanol, LC-MS/MS Mix 3 **Sexual assault**^a: dependent upon case history

SIU death investigation: Ethanol, QTOF Screen, LC-MS/MS Mix 3, IA cannabinoids

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^a dependent upon sample volume

b if fire is involved

^c other analyses may be performed dependent upon evidence/suspicion of intoxication

Appendix 4 - No Method Available

Examples of drugs/compounds for which this laboratory does not have a method

Animal toxins

α-bungarotoxin conotoxin maurotoxin tetrodotoxin

Anesthetic gases

halothane isoflurane nitrous oxide

Curare-related toxins

alloferine toxiferine tubocurarine

Other

insulin lead, mercury lithium polychlorinated biphenyls (PCB) succinylcholine thallium

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Appendix 5 - Capability of Quantitative Methods

Barbiturate method (LC-MS/MS)

amobarbital (qualitative)

butalbital pentobarbital phenobarbital phenytoin primidone secobarbital

Cannabinoid method (LC-MS/MS)

tetrahydrocannabinol (THC)
THC (11-nor-carboxy; THC-COOH)
THC (11-hydroxy; THC-OH, qualitative)
cannabidiol
cannabinol

GHB/BHB method (LC-MS/MS)

 γ -hydroxybutyrate (GHB) β -hydroxybutyrate (BHB)

LC-MS/MS Mix 2

benztropine ephedrine mitragynine (qualitative) benzylpiperazine haloperidol nicotine (semi-quantitative)

brompheniramine ketorolac pseudoephedrine

caffeine (semi-quantitative) loperamide (qualitative) trimeprazine (qualitative)

clonidine lidocaine (semi-quantitative) warfarin

LC-MS/MS Mix 3

6-monoacetylmorphine (6-MAM; diazepam (nor) methylenedioxyamphetamine qualitative) diphenhydramine methylenedioxyethylamphetamine

acetyl fentanyl etizolam methylenedioxymethamphetamine alprazolam midazolam

amitriptyline flualprazolam mirtazapine amphetamine flubromazolam morphine benzoylecgonine flunitrazepam (7-amino) nortriptyline bromazolam (qualitative) fluoxetine olanzapine oxazepam bupropion fluoxetine (nor) carfentanil flurazepam (n-desalkyl) oxycodone chlorpheniramine hydrocodone oxymorphone

chlorpheniramine hydrocodone oxymorpho citalopram/escitalopram hydromorphone paroxetine

clonazepam hydroxyrisperidone/paliperidone pseudoephedrine

clonazepam (7-amino; qualitative) isotonitazene quetiapine clonazolam ketamine risperidone clonazolam (8-amino; qualitative) ketamine (nor) sertraline cocaethylene lorazepam temazepam cocaine meperidine tramadol (cis) meperidine (nor) codeine trazodone

cyclobenzaprine mephedrone (qualitative) venlafaxine

dextromethorphanmethadonexylazinediazepammethamphetaminezopiclone

LC-MS/MS Mix 4

alprazolam (hydroxyl) doxylamine naltrexone amoxapine duloxetine nitrazepam

bromazepam flunitrazepam nitrazepam (7-amino)

buprenorphine flunitrazepam (N-desmethyl) orphenadrine

PCP butyryl fentanyl flurazepam chlordiazepoxide fluvoxamine pentazocine chlorpromazine furanyl fentanyl pheniramine clobazam imipramine promethazine levorphanol/dextrorphan clomipramine propoxyphene

(qualitative) triazolam clozapine

demoxepam loxapine triazolam (hydroxy) desipramine methylenedioxypyrovalerone trimipramine

desomorphine methotrimeprazine U-47700

methylone diltiazem venlafaxine (O-desmethyl)

diltiazem (desacetyl) methylphenidate ziprasidone doxepin naloxone zolpidem

LC-MS/MS Mix 5

acebutolol gabapentin prochlorperazine quaifenesin acetaminophen propafenone amiodarone ibuprofen propranolol amlodipine labetalol pseudoephedrine atenolol lamotrigine salicylate baclofen methocarbamol topiramate

carbamazepine (qualitative) metoprolol verapamil diflunisal naproxen vigabatrin

furosemide pregabalin

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Appendix 6 - Capability of Targeted Qualitative Methods

Urine Drug Mix (UDM; LC-MS/MS)

6-monoacetylmorphine (6-MAM)

acetyl fentanyl acetyl norfentanyl

alprazolam amitriptyline amlodipine amoxapine amphetamine

baclofen benzoylecgonine bromazepam

brompheniramine buprenorphine

buprenorphine glucuronide bupropion butyryl fentanyl

carfentanil
chlordiazepoxide
chlorpheniramine
citalopram/escitalopram

clobazam clomipramine clonazepam

clonazepam (7-amino)

clonazolam

clonazolam (8-amino)

clozapine cocaethylene cocaine

codeine

codeine-6-glucuronide cyclobenzaprine demoxepam desipramine

desomorphine dextromethorphan diazepam diazepam (nor)

diltiazem

Authorized:

diltiazem (desacetyl) diphenhydramine

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doxepin doxylamine duloxetine ephedrine etizolam fentanyl fentanyl (nor) flualprazolam flubromazolam

flunitrazepam

flunitrazepam (7-amino) flunitrazepam (N-desmethyl)

fluoxetine fluoxetine (nor) flurazepam

flurazepam (n-desalkyl) fluvoxamine

furanyl fentanyl gabapentin GHB heroin hydrocodone hydromorphone

hydromorphone-3-glucuronide

hydroxyalprazolam hydroxytriazolam imipramine ketamine ketamine (nor) lamotrigine

levorphanol/dextrorphan lidocaine

lorazepam

lorazepam glucuronide

loxapine
meperidine
meperidine (nor)
mephedrone
methadone
methamphetamine
methylenedioxyamph

methylenedioxyamphetamine methylenedioxyethylamphetamine methylenedioxymethamphetamine

methylenedioxypyrovalerone

methylone methylphenidate metoprolol midazolam mirtazapine morphine

morphine-3-glucuronide morphine-6-glucuronide

naloxone naltrexone nitrazepam

nitrazepam (7-amino)

nortriptyline olanzapine orphenadrine oxazepam

oxazepam glucuronide

oxycodone oxymorphone paroxetine pentazocine phenazepam phencyclidine pheniramine pregabalin propoxyphene propranolol pseudoephedrine

pseudoephe quetiapine risperidone sertraline tapentadol temazepam

temazepam glucuronide

THC-COOH

THC-COOH glucuronide

topiramate tramadol (cis) trazodone triazolam trimipramine U-47700 venlafaxine

venlafaxine (O-desmethyl)

zaleplon ziprasidone zolpidem zopiclone