



Patient Centered Urine Drug Testing for Primary Care

Douglas Gourlay, MD, MSc, FRCPC, FASAM

Declaration of Potential Conflict of Interest

- The content of this presentation is noncommercial and does not represent any conflict of interest



2

Learning Objectives

- Describe a patient centered approach to urine drug testing (UDT)
- Explain the differences between the clinical vs workplace/forensic test pool
- Express the importance of “testing strategy” in the clinical use of UDT
- List common myths/misconceptions of urine drug testing analysis



3

Why do we test?

- Forensics
- Treatment compliance/concurrent drug use
- Advocacy
 - With 3rd party, motivate/support behavioral change, identify abuse/addiction: Avoid “gotcha” syndrome
- Risk management

Painweek

4

Nonclinical vs Clinical Testing

- In nonclinical testing, the majority of donors are expected to be nonusers
 - Thresholds are set based on donor characteristics NOT lab capabilities
 - “-ve test results can harm the donor”
- In pain practices this is NOT the case
 - Majority of donors are user of common drugs of misuse
 - Difference is they’re legitimately +ve

Painweek

5

What’s worse than not doing UDT?

- Doing UDT inappropriately
 - Clinicians (and lab directors) must resist the urge to reach beyond the scientific AND clinical limits of the test
- Using ‘clinical’ test strategies for ‘forensic’ purposes
 - Agreeing to monitor for CPS/drug court
 - This is a dangerous practice

Painweek

6

When to Test

- Consider urine drug testing (UDT) in all patients
 - Especially those starting opioid therapy
 - When making major changes in therapy
 - In response to aberrant behavior
- Testing frequency
 - Low risk—initially and yearly if no problems?
 - High risk—weekly? Monthly if stable?
- Cheap, effective, and well tolerated by patients
 - Only patients ‘philosophically opposed’ to UDT are those patients with problems who don’t want help

Painweek

7

How to Test

- There is no ‘right or wrong’ way to test
 - Laboratory testing
 - Point-of-care (‘test strips’)
- Never do a test if you don’t know how to interpret the results
 - You must have a testing strategy
 - Need an action plan to deal with results

Painweek

8

Testing Strategies

- Routine vs random testing
 - Random more reliable, more complex
 - Routine easier to ‘prepare’ for
- Reliability
 - Witnessed require same-sex observers
 - Can use temperature strips
- How to use the results
 - Avoid “gotcha” syndrome

Painweek

9

Testing Techniques

- Presumptive
 - Immunoassay (EMIT)
- Definitive (identification)
 - GC/MS, LC/MS-MS, etc
- Point-of-care testing ('test strips')
 - Immunoassay

Painweek

10

Adulteration, Substitution, Volume Loading

- People do cheat!
- Witnessed vs unwitnessed collection
- Temperature monitoring
 - Min volume, time, within 1°C body temp
- pH, creatinine, "urine fingerprinting"
- Volume loading
 - Deliberately ingest H₂O to lower SG, Cr
 - Cr < 1.8 mmol/L AND SG < 1.003 = suspicion

Painweek

11

Drugs of Abuse

- NIDA-5 (aka "federal five")
 - Cocaine
 - Opiates
 - THC
 - Amphetamines
 - PCP
- Benzodiazepines, barbiturates, methadone, etc

Painweek

12

Cocaine

- Screen for metabolite, benzoylecgonine (BEG) NOT cocaine parent
- Metabolite $t_{1/2} \gg$ parent $t_{1/2}$
- H₂O soluble ∴ does not accumulate
- Detectable at 300ng/mL for 3-5 day
- Cocaine (parent) implies very recent drug use ie hours

Painweek

13

Opiates

- Really codeine/morphine
 - Cross reacts with many other opioids
- Threshold varies – DOT 2000 ng/mL; typically 300 ng/mL (total opioids)
- Heroin use confirmed by 6-AM (6-MAM)
 - $t_{1/2}$ short makes detection difficult
 - Never detect heroin parent*
 - Can't distinguish morphine from heroin/morphine/codeine metabolism

Painweek

14

Opioids 101

Natural	Semi-Synthetic	Synthetic
Codeine	Hydrocodone	<i>Meperidine</i>
Morphine	Hydromorphone Diacetyl Morphine	Fentanyl (Sufenta, Alfenta, Remifentanyl)
Thebaine	Oxycodone, Oxymorphone, Buprenorphine, Naltrexone, Naloxone	Methadone, Propoxyphene

Painweek

THC

- Screen looks for all cannabinoids
- Variable cut-offs (50ng/mL / 15ng/mL)
- Fat soluble
- GC/MS looks only at THC-COOH
- Infrequent users detect for <3d
- Frequent, heavy users >7d (20 ng/mL 77days positive)

Painweek

16

Amphetamines

- EMIT screens triggered with decongestants, antihistamines
- May react to MDMA (Ecstasy), MDA etc
- Many prescription and OTC drugs give false positive EMIT screens
- Cut-off 1000 ng/mL, confirms 500 ng/mL
- "Vicks Nasal Inhaler" dilemma (USA)
- Typically detectable for <3 d

Painweek

17

PCP

- Phencyclidine (also reacts with Ketamine)
- Low yield except with specific patient populations *in certain areas*
- Cut-off of 25 ng/mL
- Detectable for < 7 d

Painweek

18

Other Drugs

- Specific opioids
 - Hydromorphone—may need to ask lab for assistance
 - Oxycodone—needs specific assay
 - Hydrocodone
 - Buprenorphine (immunoassay)
 - Methadone/fentanyl do NOT yield +ve ‘opiate’ screens
- Benzodiazepines
 - Difficult to reliably detect, especially clonazepam even when abused—check with lab regarding sensitivity

Painweek

19

Poppy Seeds

- Poppy seeds don't give false positives
 - They lead to TRUE positives
 - Can exceed DOT cutoffs for several hours
 - May show both morphine and codeine
 - NEVER accounts for 6-MAM
- *People on UDT programs should not eat poppy seeds*

Painweek

20

Passive Marijuana

- ‘Incidental’ exposure does not lead to +ve UDT
- Depends on cut off concentration
- Can not easily distinguish prescribed oral THC from smoked marijuana
- Single use does not lead to persistent +ve results

Painweek

21

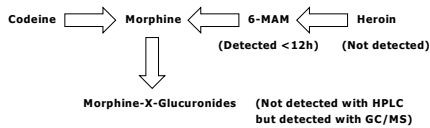
Passive Cocaine

- Nasal cocaine (cocaine HCl) can not be put in cigarette to give positive result
 - Crack cocaine can lead to positive result
- Cocaine base sublimates when heated ∴
 - Found on many surfaces where crack cocaine is used
 - \$20 bills frequently test positive for cocaine
- Medical uses result in positive results
 - ENT, ophthalmology, plastic surgery

Painweek

22

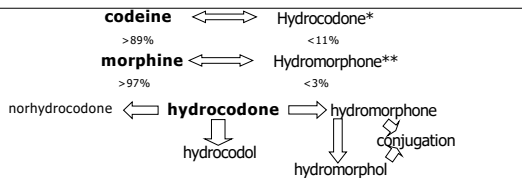
Opioid Metabolism



Painweek

23

Drug Testing Traps



Baselt RC. *Disposition of Toxic Drugs and Chemicals in Man*. 7th ed 2004

* Identification of Hydrocodone in Human Urine Following Controlled Codeine Administration, JM Oyler et al *Journal of Analytical Toxicology* 24(7) 2009 p530-535

** Evidence of Morphine Metabolism to hydromorphone in pain patients chronically treated with morphine, E Cone et al, *Journal of Analytical Toxicology* 30(1) 2006 p1-5

Painweek

24

What to do with unexpected results?

- First, call the lab
 - If unexpected +ve, check for legitimate reasons for true positives
 - ENT cocaine for epistaxis
 - Morphine in codeine user
 - Hydrocodone in codeine user (~11% or less)
 - If unexpected –ve, check for test sensitivity, subthreshold results, dilute sample, lab error
- Speak with patient
 - Ask about ALL drug use including OTC and time of last use
 - When truly negative, look for bingeing (ie, running out)
- Never ignore an aberrant result!

Painweek

27

Test Interpretation Traps: Urinary Levels

- Urinary drug and drug metabolite excretion are a function of many factors which may not be static
 - Volumes of distribution, urinary pH, state of hydration, time of last dose, GI absorption effects etc
 - It is unwise to draw any conclusions based solely or largely on urinary analyte concentrations
 - Drug testing is the beginning not the end of discussion— use 'social engineering' to solve the problem

Painweek

28

Using the Results: (it's all in the strategy)

- First, do the results "fit" ?
 - If yes, could they be 'hiding' an abnormal result? ie, +ve opiates / +ve bzd
 - Beware of the expected analyte
- Compliance testing
 - What does the –ve mean?
 - Have a diff Dx for the unexpected result
- Can you interpret the results?
 - Ask before collecting sample
 - New meds? New OTC drugs? Recreational use?

Painweek

29

Approaching the Patient

- “Offer” drug testing to the patient
 - Majority of patients will have no problems with UDT
 - If patient is ‘philosophically opposed’ to UDT, bodes poorly for this patient
 - Remind patient that this will severely limit the pharmacologic choices for treating their pain
 - Reassure the patient that UDT is part of a comprehensive risk management strategy

Conclusions

- UDT, when done with respect and sensitivity can be an important part of a comprehensive care plan for all, not only high risk patients that
 - Reduces patient stigma
 - Improves patient care
 - And hopefully, reduces risk
- The clinical context is essential for proper UDT interpretation
 - Risk management is FOR the patient

Resources

- <http://www.UDTmonograph6.com>
 - For UDT monograph
- dgourlay@cogeco.ca

References

1. D Gourlay, HA Heit (co-authors), Y Caplan: Urine Drug Testing in Clinical Practice, The Art and Science of Patient Care. <http://www.udtmonographs.com/> 6th Edition, August 31, 2015.
2. HA Heit, D L Gourlay: Urine Drug Testing in Pain Medicine: J Pain Sympt. Manage. 2004;27(3): 260-67
3. E.J. Cone, H.A. Heit, Y.H. Caplan, D. Gourlay; J. Anal. Toxicol.: Evidence of Morphine Metabolism to Hydromorphone in Pain Patients Chronically Treated with Morphine, 2006;30(1):1-5.
4. Sloan PA, Barkin RL: Oxymorphone and Oxymorphone Extended Release: A Pharmacotherapeutic Review J of Opioid Management 4(3), May/June 2006; 131-44.
5. Natziger AN et al. Utility and Application of UDT in Chronic Pain Management With Opioids Clin J Pain. 2009;25(9):73-79
6. DL Gourlay, HA Heit: The Art and Science of Urine Drug Testing. Clin J Pain. 2010;26(4):358.
7. MROALERT, November 6, 2006: Vol.XVII; No. 9(1-4)
8. DL Gourlay, HA Heit. Urine Drug Testing in Pain and Addiction Medicine. In H Smith and SD Passik (eds) Pain and Chemical Dependency. New York: Oxford University Press, 2008: 353-58.
9. DL Gourlay, HA Heit. Compliance Monitoring in Chronic Pain Management. In S M Fishman, JC Ballantyne, JP Rathmell, (eds). Bonica's Management of Pain, Fourth Edition. Philadelphia: Lippincott Williams & Wilkins, 2010: 854-859. 5th edition in press
10. Katz, N., Fanciullo, G.J. The Clinical J of Pain, Vol. 18, No. 4, S 2002: S76-82

Painweek
