

# Patient Centered Urine Drug Testing for Primary Care

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# **Declaration of Potential Conflict of Interest**

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

2

3

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# 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

## Why do we test?

Forensics

Treatment compliance/concurrent drug use

Advocacy

-With 3<sup>rd</sup> party, motivate/support behavioral change, identify abuse/addiction: Avoid "gotcha" syndrome

4

5

6

Risk management

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# **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
  - $-\ensuremath{``\text{-ve}}$  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

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# 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

# 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

7

8

9

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# 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

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# **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

# **Testing Techniques**

Presumptive

-Immunoassay (EMIT)

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

-Immunoassay

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## Adulteration, Substitution, Volume Loading

10

11

- 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 H<sub>2</sub>O to lower SG, Cr • Cr<1.8 mmol/L AND SG<1.003 = suspicion

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# **Drugs of Abuse**

NIDA-5 (aka "federal five")

- -Cocaine
- -Opiates
- -THC
- -Amphetamines
- -PCP

Benzodiazepines, barbiturates, methadone, etc

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12

## Cocaine

- Screen for metabolite, benzoylecgonine (BEG) NOT cocaine parent
- Metabolite t ½ >> parent t ½
- $\bullet \, H_2O \text{ soluble } \therefore \text{does not accumulate}$
- Detectable at 300ng/mL for 3-5 day
- Cocaine (parent) implies very recent drug use ie hours

13

14

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# **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)
  - $-t_{\frac{1}{2}}$  short makes detection difficult
  - -Never detect heroin parent\*
  - -Can't distinguish morphine from heroin/morphine/codeine metabolism

Natural	Semi- Synthetic	Synthetic
Codeine	Hydrocodone	Meperidine
Morphine	Hydromorphone Diacetyl Morphine	Fentanyl (Sufenta, Alfenta, Remifentanil
Thebaine	Oxycodone, Oxymorphone, Buprenorphine. Naltrexone, Naloxone	Methadone, Propoxyphene



# THC

Screen looks for all canabinoids

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

16

17

18

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# 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</p>

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# 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</li>

# **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

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# **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

20

21

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# **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

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7

## **Passive Cocaine**

 Nasal cocaine (cocaine HCI) can not be put in cigarette to give positive result

22

- -Crack cocaine can lead to positive result
- $\hfill \black$  Cocaine base sublimates when heated  $\hfill \hfill \black$ 
  - $-\ensuremath{\mathsf{Found}}$  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







## 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

27

28

29

- 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!

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# **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 discussionuse 'social engineering' to solve the problem

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#### 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?

# **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

30

32

33

 Reassure the patient that UDT is part of a comprehensive risk management strategy

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## 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

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### **Resources**

http://www.UDTmonograph6.com

-For UDT monograph

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