Influence of Five Adulterants on Detection and Quantification of Tramadol in Urine Samples

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Abstract

Background: There is a growing evidence of abuse of tramadol in some African and West Asian countries considering large seizures of such preparations in North and West Africa especially in Egypt. Urine testing for drugs of abuse has become an integral weapon in the nation’s war against drugs. A limitation inherent in all urine drug testing is the possibility of sample adulteration or substitution. Aim of study: To detect qualitative and quantitative effects of five adulterants on positive urine samples for tramadol. Subject and Method(s): This study was conducted in Clinical Toxicology Laboratory in Sohag University Hospitals. The samples were tested for its integrity by checking PH, specific gravity and creatinine. The samples were tested by RIA then confirmed and quantified by HPLC. Results: Urine samples adulterated with vinegar, drano and liquid hand soap generated false negative results by immunoassay testing. HPLC confirmation showed decrease tramadol conc. below limit of quantification in urine samples adulterated with 40% vinegar and 40% drano. Conclusion: Some adulterants make it easy to produce false negative results and the specimen integrity testing is inadequate in detection of these adulterants.

Key words
Drug abuse, Tramadol detection, Urine adulteration, HPLC confirmation

Introduction

Substance abuse in Egypt is a serious public health threat. Recent studies have demonstrated increase in the prevalence of the use of cannabis and tramadol (Saleh, 2015).

Tramadol is a centrally acting analgesic with a multimode of action. It acts on serotonergic and noradrenergic nociception, while its metabolite O-desmethyl tramadol acts on the μ-opioid receptor (Babalonis et al., 2013).

Tramadol is generally considered as a medicinal drug with a low potential for dependence relative to morphine. Nevertheless, tramadol dependence may occur when used for prolonged periods of time (more than several weeks to months). Dependence to tramadol may occur when used within the recommended dose range of tramadol but especially when used at supra-therapeutic doses (Alvarado et al., 2005).

There is growing evidence of abuse of tramadol in some African and West Asian countries considering large seizures of such preparations in North and West Africa. Abuse of tramadol is reported by Egypt, Gaza, Jordan, Lebanon, Libya, Mauritius, Saudi Arabia and Ghana (Ellison et al., 2018).

According to Nabil et al., (2015) the prevalence of tramadol dependency according to all substance abusers was 49%. The prevalence of comorbid psychiatric disorders was 43%.

Urine testing for drugs of abuse has become an integral weapon in the nation’s war against drugs. These drug tests are used in pre-employment screening, post-accident assessment, probation control, and inmate determent (Wong, 2002).

Drug testing occurs in two phases: screening and confirmation. Historically, the initial screen has been a chemical test or immunoassay (IA) that can provide a reasonable turnaround time with minimal labor and resources (Al-khayal et al., 2017).

A limitation inherent in all urine drug testing is the possibility of sample adulteration or substitution (Jaffee et al., 2007).

Adulteration process is defined as the tampering or manipulation of a urine sample with the intention of changing the test results. Urine sample adulteration is very serious problem in forensic urine drug testing process. Sample adulteration is usually performed by substitution, dilution or the addition of adulterants agents including so called "masking agents" sold commercially (Ragab et al., 2018).

The use of adulterant agents can cause false negative results in drug tests by either interfering with the screening test procedure and/or destroying the drugs present in the urine sample (Yee et al., 2014).
**Aim of the Work**

To detect qualitative and quantitative effects of five adulterants on positive urine samples for tramadol.

**Subjects and Methods**

This study was conducted in Clinical Toxicology Laboratory in Sohag University Hospitals. Acceptable samples had the following criteria: from 10-100ml urine in volume, voided in a clean dry labeled container without preservative. The samples were tested by immunoassay for tramadol, positive samples only were included. Urine samples were tested for integrity including measurement of (PH, specific gravity, creatinine) before testing.

Five Types of adulterants were used.

- Vinegar, bleach, visine eye drops and liquid drano at 3 concentrations 10%, 20% and 40%.
- Liquid hand soap at 3 conc. 5%, 10% and 20%.

These adulterants levels were selected to obtain an accurate representation of real-world samples adulteration as it would easy to be brought in a small container and added to the urine samples.

- To prepare adulterated samples, 1 mL aliquots were obtained from the urine samples for each drug. 1 ml of the unadulterated urine sample was used to determine the initial concentration of the drug by High Performance Liquid Chromatography (HPLC).

The total volume of the adulterated samples was maintained at 1ml. The amount of liquid adulterants was added to the urine sample to reach the 1ml limit. The 10 % v/v sample had 900 μL of urine and 100 μL of adulterant. This process was followed for the remaining concentrations: 20 % v/v (800:200) 40 % v/v (600:400) and 5%v/v (950:50).

Screening of tramadol in urine samples by immunoassay:

Apparatus: Radio immunoassay apparatus using drug analyzer: (CDX90), Thermofisher Scientific co. supplier AMG Company.

Fully automated random access analyzer, dedicated drug testing system (photometric).

Serial number 7218-0150 present in Clinical Toxicology Laboratory in Sohag University Hospital.

i- Principle of procedure:

This assay is a semi-quantitative assay based on the competition of Tramadol labeled enzyme glucose -6-phosphate dehydrogenase (G6PDH) and the free drug in the urine sample for the fixed amount of antibody binding sites.

In the absence of the free drug in the sample, the antibody binds the drug enzyme conjugate and enzyme activity is inhibited. The enzyme (G6PDH) activity is determined at 340nm spectrophotometrically by the conversion of NAD to NADH

ii- Calibration: Figure (A)

1- Construct the calibration curve (log-logit mode) by using the following calibrators:

- Negative Urine Calibrator.
- 100 ng/ml Urine Calibrator.
- 200 ng/ml Urine Calibrator (Cutoff calibrator).
- 500 ng/ml Urine Calibrator.
- 1000 ng/ml Urine Calibrator.

2- For qualitative analysis use the 200 ng/ml calibrator as a cutoff level to distinguish “positive” and “negative” specimens.

3- Check the constructed calibration curve using the provided QC materials 150, 250ng/ml as 150 ng/ml considered LQC which control negative results and HQC250 ng/ml control positive results.

Each conc. of each adulterant was added separately to urine sample and the samples were examined for specimen integrity then retested by immunoassay where cut off for tramadol detection was 200 ng/ml above it considered positive and below it considered negative and finally the samples were confirmed by HPLC.

Confirmation and quantification of tramadol in urine samples by HPLC:

Apparatus: High Performance Liquid Chromatography (HPLC) (Agilent; USA): Consisted of an Agilent technologies 1200 series quaternary pump combined with an Agilent 1200 series photo diode array detector (USA), an Agilent 1200 series vacuum degasser (USA) and an Agilent autosampler injector.

Chromatographic separation was performed on a Zorbax SB C8 (250 mm x4.6 mm, 5 μm) column (USA). HPLC present in Sohag Clinical Toxicology Lab– Sohag University Hospitals.

i- Calibrators and quality control

A stock solution of tramadol (T), o-desmethyl tramadol (ODT) and propranolol used as internal standard (IS) were prepared at a concentration of 1 mg/mL in methanol and kept stored at -20C. Intermediate standards at concentration. of 100 mg/mL for each analyte were prepared in methanol by diluting from 1.0 mg/mL stock standards. Different stock standards were used to. Prepare quality control samples (QCs) at the same concentrations. Working calibrators (250, 500, 750, 1000, 1500 and 2000 ng/mL) of T and ODT were made by a serial dilution of the intermediate solution with drug free human urine. QCs were prepared from a separate stock solution at concentrations of 450, 900 and 1800 ng/mL. A working standard solution of 5.00 Ug/mL propranolol (IS) was prepared by diluting propranolol stock solution with distilled water.

ii- Phosphate buffer preparation:

Phosphate buffer (0.01 M) was prepared by dissolving 1.36 g (0.01 mol) of potassium dihydrogen phosphate in 1 L deionized water.

iii- Extraction procedure

To 10-mL polypropylene tubes added 1.0 mL of urine, 75 mL of 5 Ug/mL propranolol (IS), 100 mL of conc. ammonium hydroxide (33%) and 6.0 mL of MTBE. The tubes were then mixed by rotator at the rate of 40 rpm for 20-min and centrifuged at 3200 rpm for 5-min. The organic layer was transferred to 10-mL polypropylene tube containing 0.5 mL of 1.0 M hydrochloric acid. The
tubes were then vortex mixed for 5-min and centrifuged at 3200 rpm for 5-min. The organic layer was discarded. To the remaining aqueous solution, 150 mL of conc. ammonium hydroxide and 2.0 mL of MTBE were added. The tubes were then centrifuged at 3200 rpm for 5-min. The organic layer was transferred to 5-mL glass tubes and evaporated to dryness. The dried extracts were reconstituted in 200 mL acetonitrile, vortex mixed for 30-s and 100 mL was injected into the HPLC system.

- Check PH at every step and adjust it according to result.

iv- HPLC analysis:
Chromatography condition:

   The column oven temperature was maintained at 25 °C. The mobile phase consisted of ACN: phosphate buffer (40:60, v/v) at a flow rate of 0.5 mL/min. The detector was set to scan from 200 to 360 nm and had a discrete channel set at 218 nm, which was the wavelength used for quantification.

Calibration curve construction: Figure (B)

   Linearity of the method was investigated by evaluation of the regression line and expresses by coefficient of determination (r2). Linearity was achieved with a minimal r2 of 0.993. Calibration curves were prepared by spiking blank urine with corresponding analytical working solutions to obtain calibration concentrations within 250–2000 ng/mL. Negative QCs were analyzed after each linearity sample to evaluate potential carry-over.

Results

Specimen integrity test:

A. Effect of vinegar on specimen integrity tests
   - Addition of 40 % vinegar to urine samples lead to decrease in PH and creatinine below their normal range while specific gravity increased.
   - Addition of 20% vinegar also lead to decrease in PH and increase in specific gravity but creatinine level was not affected.
   - Finally addition of 10 % vinegar has no effect on PH, specific gravity, or creatinine.

B. Effect of bleach on specimen integrity tests
   - Addition of 40 % bleach to urine samples lead to increase in PH and specific gravity above their normal range while creatinine decreased.
   - Addition of 20% bleach also lead to increase in PH and specific gravity but creatinine level was not affected.
   - Finally addition of 10% bleach has no effect on PH, specific gravity, or creatinine.

C. Effect of visine on specimen integrity tests
   - Addition of visine to urine samples has no effect on PH, specific gravity, or creatinine.

D. Effect of drano on specimen integrity tests
   - Addition of 40 % drano to urine samples lead to increase in PH above normal range while specific gravity and creatinine decreased.
   - Addition of 20% drano also lead to increase in PH but specific gravity and creatinine decreased.
   - Finally addition of 10 % drano has no effect on PH, specific gravity, or creatinine.

E. Effect of liquid hand soap on specimen integrity tests
   - Addition of 20% liquid hand soap to urine samples lead to increase in PH and specific gravity above their normal range while creatinine level decreased.
   - Addition of 10% liquid hand soap also leads to increase in PH and specific gravity while creatinine level not affected.
   - Finally addition of 5% liquid hand soap has no effect on PH or creatinine and increase in specific gravity.

Imunoassay screening for tramadol:

1. The parent sample concentration was 530 ng/ml. (as the method of detection considered semiquantitative not only screening).
   - Addition of vinegar at high conc. 40 % is able to successfully masking positive response of tramadol in tested urine samples. While moderate conc.20 % and low conc.10% cannot affect tramadol detection in urine samples as shown in table (1) and figure (1).

2. On other hand addition of bleach, visine whatever their conc. failed to mask tramadol detection by immunoassay as shown in table (1) and figure (1).

3. For drano it was effective for decreasing the response rate for tramadol using immunoassay method at high conc. 40% and moderate 20%. While 10% is still has no effect on tramadol result.as shown in table (1) and figure (1).

4. Unfortunately, addition of liquid hand soap by any conc. even low conc. up to5% can mask tramadol detection by immunoassay giving false negative results.as shown in table (2) and figure (1).

5. Detection and quantification tramadol and O-des methyl tramadol (ODT) by HPLC:
   - The parent positive sample conc. was: Tramadol 494ng/ml and ODT 804 ng/ml Figure (4).
   - Limit of detection (LOD): 150 ng/ml
   - Limit of quantification (LOQ): 250 ng/ml
   - 150ng/ml-250ng/ml: The drug can be detected but cannot be quantified.

1- Effect of vinegar on Tramadol and ODT detection & quantification by HPLC
   - After addition of vinegar in conc. 40% leading to decrease in conc. of tramadol to less than LOQ but still be detectable (192.2 ng/ml). For ODT also decrease to more or less half the actual conc. (479.5ng/ml).
   - While addition of 20% vinegar decrease tramadol conc. more or less half the actual conc. (257.5 ng/ml). For ODT decrease to lesser degree (636.4).
   - Finally 10% has the least effect as there is minimal decrease from actual conc. (439.7ng/ml) and more or less the same occurred in ODT conc. (690.5ng/ml).As shown in table (3) and figures (5,6&7).
2. Effect of bleach on Tramadol and ODT detection & quantification by HPLC
   - After Addition of bleach in conc.40% decrease tramadol conc. to (300.5ng/ml). For ODT also decrease to more or less half the actual conc.(465.2ng/ml).
   - While addition of 20% bleach decrease tramadol conc. to (392.5 ng/ml) and for ODT decrease to less degree (637.4ng/ml).
   - Finally 10% has the least effect as there is minimal decrease from actual conc. (437.6ng/ml) and more or less the same occurred in ODT conc. (656.5ng/ml). As shown in table (3) and Figures (8,9&10).

3. Effect of visine on Tramadol and ODT detection & quantification by HPLC
   - After Addition of visine in conc.40% decrease tramadol conc. to (301.8ng/ml).and for ODT also decrease to more or less half the actual conc. (474.3 ng/ml).
   - While addition of 20% visine decrease tramadol conc. to (389.7ng/ml) and for ODT decrease to less degree (631.5ng/ml).
   - Finally 10% has the least effect as there is minimal decrease from actual conc. (451.2ng/ml) and more or less the same occurred in ODT conc. (677.2ng/ml). As shown in table (3) and figures (11,12&13).

4. Effect of drano on Tramadol and ODT detection & quantification by HPLC
   - Addition of drano in conc.40% leading to decrease in conc. of tramadol to less than LOQ but still be detectable (231.2 ng/ml).and for ODT also decrease to more than half the actual conc.(332.1ng/ml).
   - While addition of 20% drano cause moderate decrease in tramadol conc (398.4 ng/ml) and for ODT decrease to less degree (645.2 ng/ml).
   - Finally 10% has the least effect as there is minimal decrease from actual conc.(433.1ng/ml) and more or less the same occurred in ODT conc. (685.4ng/ml) as shown in table (3) and figures (14,15 & 16).

5. Effect of liquid hand soap on Tramadol and ODT detection & quantification by HPLC
   - Addition of liquid hand soap in conc.20% leading to moderate decrease in conc. of tramadol (323.6 ng/ml) and marked decrease in ODT conc. (359.1ng/ml).
   - While addition of 10% liquid hand soap cause decrease in tramadol conc. to (414.9 ng/ml) and for ODT moderate decrease to (439.6 ng/ml).
   - Finally 5% has the least effect on tramadol as there is minimal decrease from actual conc. (432.4ng/ml) and moderate decrease in ODT conc. (507.7ng/ml).As shown in table (4) and figures (17, 18&19).

Statistical study for influence of different adulterants on Tramadol and ODT quantification by HPLC showing that addition of vinegar, drano and liquid hand soap decrease tramadol and ODT significantly with increased concentration as shown in table (5).

<table>
<thead>
<tr>
<th>Adulterant Conc.</th>
<th>Vinegar</th>
<th>Bleach</th>
<th>Visine</th>
<th>Drano</th>
</tr>
</thead>
<tbody>
<tr>
<td>40%</td>
<td>153ng/ml</td>
<td>320ng/ml</td>
<td>351ng/ml</td>
<td>91ng/ml</td>
</tr>
<tr>
<td></td>
<td>(Negative)</td>
<td>(Positive)</td>
<td>(Positive)</td>
<td>(Negative)</td>
</tr>
<tr>
<td>20%</td>
<td>363ng/ml</td>
<td>401ng/ml</td>
<td>409ng/ml</td>
<td>133ng/ml</td>
</tr>
<tr>
<td></td>
<td>(Positive)</td>
<td>(Positive)</td>
<td>(Positive)</td>
<td>(Negative)</td>
</tr>
<tr>
<td>10%</td>
<td>415ng/ml</td>
<td>451ng/ml</td>
<td>459ng/ml</td>
<td>251ng/ml</td>
</tr>
<tr>
<td></td>
<td>(Positive)</td>
<td>(Positive)</td>
<td>(Positive)</td>
<td>(Positive)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Liquid hand soap conc.</th>
<th>Tramadol</th>
</tr>
</thead>
<tbody>
<tr>
<td>20%</td>
<td>Zero</td>
</tr>
<tr>
<td></td>
<td>Negative</td>
</tr>
<tr>
<td>10%</td>
<td>53ng/ml</td>
</tr>
<tr>
<td></td>
<td>Negative</td>
</tr>
<tr>
<td>5%</td>
<td>60ng/ml</td>
</tr>
<tr>
<td></td>
<td>Negative</td>
</tr>
</tbody>
</table>
Table (3): Effect of different adulterants conc. on tramadol and o-des methyl tramadol (ODT) detection and quantification by HPLC

<table>
<thead>
<tr>
<th>Conc./adulterant</th>
<th>Vinegar</th>
<th>Bleach</th>
<th>Visine</th>
<th>Drano</th>
</tr>
</thead>
<tbody>
<tr>
<td>40%</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tramadol</td>
<td>190</td>
<td>-ve</td>
<td>300.5</td>
<td>+ve</td>
</tr>
<tr>
<td>ODT</td>
<td>479.5</td>
<td>+ve</td>
<td>465.2</td>
<td>+ve</td>
</tr>
<tr>
<td>20%</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tramadol</td>
<td>257.5</td>
<td>+ve</td>
<td>392.5</td>
<td>+ve</td>
</tr>
<tr>
<td>ODT</td>
<td>636.4</td>
<td>+ve</td>
<td>637.4</td>
<td>+ve</td>
</tr>
<tr>
<td>10%</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tramadol</td>
<td>439.7</td>
<td>+ve</td>
<td>437.6</td>
<td>+ve</td>
</tr>
<tr>
<td>ODT</td>
<td>690.5</td>
<td>+ve</td>
<td>656.5</td>
<td>+ve</td>
</tr>
</tbody>
</table>

Table (4): Effect of liquid hand soap conc. on tramadol and o-des methyl tramadol (ODT) detection and quantification by HPLC

<table>
<thead>
<tr>
<th>Conc./adulterant</th>
<th>Tramadol</th>
<th>ODT</th>
</tr>
</thead>
<tbody>
<tr>
<td>20%</td>
<td>323.6</td>
<td>+ve</td>
</tr>
<tr>
<td>10%</td>
<td>414.9</td>
<td>+ve</td>
</tr>
<tr>
<td>5%</td>
<td>432.4</td>
<td>+ve</td>
</tr>
</tbody>
</table>

Table (5): Statistical study for influence of adulternants on tramadol and ODT quantification by HPLC.

<table>
<thead>
<tr>
<th>Adulterant</th>
<th>Sample size (N)</th>
<th>Pearson correlation (tramadol)</th>
<th>Pearson correlation (ODT)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vinegar</td>
<td>12</td>
<td>-0.972</td>
<td>-0.994</td>
</tr>
<tr>
<td>Bleach</td>
<td>12</td>
<td>-0.669</td>
<td>-0.743</td>
</tr>
<tr>
<td>Visine</td>
<td>12</td>
<td>-0.721</td>
<td>-0.811</td>
</tr>
<tr>
<td>Drano</td>
<td>12</td>
<td>-0.997</td>
<td>-0.895</td>
</tr>
<tr>
<td>Liquid hand soap</td>
<td>12</td>
<td>-0.851</td>
<td>-0.876</td>
</tr>
</tbody>
</table>

Significance at 1 % level

Figure (1): Effects of different adulterants on tramadol detection by RIA
**HPLC Figures**

Figure (2): Chromatogram for negative quality control (blank urine)

Figure (3): Chromatogram for one calibrator 1000ng/ml

Figure (4): Chromatogram for the parent sample (Tramadol conc. 494ng/ml ODT conc. 804ng/ml)

Figure (5): Chromatogram for parent sample after addition of 40% vinegar (Tramadol conc. 192.2ng/ml & ODT conc. 479.5ng/ml)

Figure (6): Chromatogram for parent sample after addition of 20% vinegar (Tramadol conc. 257.5ng/ml & ODT conc. 636.4ng/ml)

Figure (7): Chromatogram for parent sample after addition of 10% vinegar (Tramadol conc. 439.7ng/ml & ODT conc. 690.5ng/ml)
Figure (8): Chromatogram for parent sample after addition of 40% bleach (Tramadol conc. 300.5 ng/ml & ODT conc. 465.2 ng/ml)

Figure (9): Chromatogram for parent sample after addition of 20% bleach (Tramadol conc. 392.5 ng/ml & ODT conc. 637.4 ng/ml)

Figure (10): Chromatogram for parent sample after addition of 10% bleach (Tramadol conc. 437.6 ng/ml & ODT conc. 656.5 ng/ml)

Figure (11): Chromatogram for parent sample after addition of 40% visine (Tramadol conc. 301.8 ng/ml & ODT conc. 474.3 ng/ml)

Figure (12): Chromatogram for parent sample after addition of 20% visine (Tramadol conc. 389.7 ng/ml & ODT conc. 631.5 ng/ml)

Figure (13): Chromatogram for parent sample after addition of 10% visine (Tramadol conc. 451.2 ng/ml & ODT conc. 677.2 ng/ml)
Figure (14): Chromatogram for parent sample after addition of 40% drano (Tramadol conc. 231.2 ng/ml & ODT conc. 332.1 ng/ml)

Figure (15): Chromatogram for parent sample after addition of 20% drano (Tramadol conc. 398.4 ng/ml & ODT conc. 645.2 ng/ml)

Figure (16): Chromatogram for parent sample after addition of 10% drano (Tramadol conc. 433.1 ng/ml & ODT conc. 685.4 ng/ml)

Figure (17): Chromatogram for parent sample after addition of 20% liquid hand soap (Tramadol conc. 323.6 ng/ml & ODT conc. 359.1 ng/ml)

Figure (18): Chromatogram for parent sample after addition of 10% liquid hand soap (Tramadol conc. 414.9 ng/ml & ODT conc. 439.6 ng/ml)

Figure (19): Chromatogram for parent sample after addition of 5% liquid hand soap (Tramadol conc. 432.4 ng/ml & ODT conc. 507.7 ng/ml)
Figure (A): Tramadol calibration curve on radioimmunoassay.

Figure (B): Tramadol calibration curve on HPLC

Discussion

Regarding effect of vinegar on specimen integrity tests: addition of 40% vinegar to urine samples lead to decrease in PH and creatinine below their normal range while specific gravity increased, addition of 20% vinegar also lead to decrease in PH and increase in specific gravity but creatinine level not affected and, finally addition of 10% vinegar has no effect on PH, specific gravity, nor creatinine.

The above result is in consistent with Olivieri et al., (2018) who found that vinegar appears to lower pH levels, which can affect binding, reaction times, and drug solubility.

Considering radio immunoassay screening for tramadol in this study: addition of vinegar at high conc. 40% v/v is able to successfully masking positive response of tramadol in tested urine samples, while moderate conc. 20% and low conc. 10% cannot affect tramadol detection in urine samples.

However HPLC reveals that addition of vinegar in conc. 40% leading to decrease in conc. of tramadol to less than LOQ but still be detectable (192.2 ng/ml), and for ODT also decrease to more or less half the actual conc. (479.5ng/ml), while addition of 20% vinegar decrease tramadol conc. more or less half the actual conc.(257.5 ng/ml) and for ODT decrease to less degree (636.4). Finally addition of 10% has the least effect as there is minimal decrease from actual conc. Of tramadol (439.7ng/ml) and more or less the same occurred in ODT conc(690.5ng/ml).

This was in agreement with Paul et al., (2000) who noticed considerable decrease in free opioids at a lower PH. In other research, Thabet et al., (2016) reported increase in the acidity causes highly significant reduction in the drug level.

Regarding effect of bleach on specimen integrity tests: addition of 40% bleach to urine samples lead to increase in PH and specific gravity above their
normal range while creatinine decreased. Addition of 20% bleach also lead to increase in PH and specific gravity but creatinine level not affected and finally, addition of 10 % bleach has no effect on PH, specific gravity, or creatinine. On other hand addition of bleach whatever its conc. failed to mask tramadol detection by immunoassay.

However HPLC reveals that addition of bleach in conc.40% decrease tramadol conc. to (300.5ng/ml),and for ODT also decrease to more or less half the actual conc. (465.2ng/ml), addition of 20% bleach decrease tramadol conc. to (392.5 ng/ml) and for ODT decrease to less degree (637.4ng/ml). Finally 10% has the least effect as there is minimal decrease from actual conc. (437.6ng/ml) and more or less the same occurred in ODT conc. (656.5ng/ml).

In contrary to the above results, olivieri et al., (2018) stated that bleach is extremely effective in adulterating urine screens positive for benzodiazepines, cocaine, THC and opiates in the EIA.

This is in consistent with Thabet et al., (2016) who reported increase in alkalinity causes apparent increase in drug level in case of high tramadol concentration.

Furthermore, addition of visine to urine samples has no effect on PH, specific gravity, nor creatinine, and these results matching with El Khateeb and Arafa, (2019).

On other hand in contrary to El Khateeb and Arafa, (2019) addition of visine eye drops failed to mask tramadol detection by immunoassay. Addition of visine in conc.40% decrease tramadol conc. to (301.8ng/ml), and for ODT also decrease to more or less half the actual conc. (474.3ng/ml). While addition of 20% visine decreases tramadol conc. (389.7ng/ml), and for ODT decrease to less degree (631.5ng/ml). Finally 10% has the least effect as there is minimal decrease from actual conc. (451.2ng/ml) and more or less the same occurred in ODT conc. (677.2ng/ml).

The active ingredient in Visine eye drops is tetrahydro-zoline hydrochloride, which relieves redness and irritation by constricting blood vessels. However, Dasgupta, (2007) reported that the mechanism of adulteration is most likely due to the inactive ingredients benzalkonium chloride and borate. Visine is effective at masking THC metabolites, but not other drugs, across various immunoassays.

Regarding effect of drano on specimen integrity tests: addition of 40 % drano to urine samples lead to increase in PH above normal range while specific gravity and creatinine decreased. Also, addition of 20% drano also lead to increase in PH but specific gravity and creatinine decreased. Finally addition of 10 % drano has no effect on PH, specific gravity, nor creatinine. In consistent with Fu et al., (2014) Drano causes change to alkaline pH in urine samples, which may affect reaction rates, drug solubility, and binding. It is also effective for decreasing the response rate for tramadol using immunoassay method at high conc.

40% and moderate conc. 20% while 10% is still has no effect on tramadol result.

However, HPLC found that addition of drano in conc.40% leading to decrease in conc. of tramadol to less than LOQ but still be detectable (231.2 ng/ml).and for ODT also decrease to more than half the actual conc. (332.1ng/ml).

While addition of 20% drano causes moderate decrease in tramadol conc. (398.4 ng/ml) and for ODT decrease to less degree (645.2 ng/ml).

Finally 10% has the least effect as there is minimal decrease from actual conc. (433.1ng/ml) and more or less the same occurred in ODT conc. (685.4ng/ml).

In consistent with Oliveri et al., (2018) who stated that drano has been reported to cause strong adulterating effects on EIA. Also he suggest that Drano’s oxidation reaction to the drug assays is the primary mechanism of adulteration.

For effect of liquid hand soap on specimen integrity tests, in consistent with Dasgupta (2010) who reported that soap may alter pH levels in urine samples and may also interfere with drug binding on immunoassay. in the present study it was found that addition of 20 % liquid hand soap to urine samples lead to increase in PH and specific gravity above their normal range while creatinine level decreased. Addition of 10% liquid hand soap also lead to increase in PH and specific gravity while creatinine level not affected and finally addition of 5% liquid hand soap has no effect on PH nor creatinine and increase in specific gravity.

Unfortunately, in consistent with Wu (2003) who stated that dishwashing detergent adulteration has caused false-negative results across a variety of drug assays using the CEDIA, including screens for many drugs including tramadol (amphetamine, barbiturates, cocaine, opiates, PCP, and THC) we also found that addition of liquid hand soap by any conc. even low conc. up to 5% can mask tramadol detection by immunoassay giving false negative results.

However, HPLC results reveals that addition of liquid hand soap in conc.20% leading to moderate decrease in conc. of tramadol (323.6 ng/ml) and marked decrease in ODT conc.(359.1ng/ml), while addition of 10% liquid hand soap cause decrease in tramadol conc. to (414.9 ng/ml) and for ODT moderate decrease to (439.6 ng/ml). Finally 5% has the least effect on tramadol as there is minimal decrease from actual conc. (432.4ng/ml) and moderate decrease in ODT conc. (507.7ng/ml).

Most detergents and soaps contain multiple ingredients including surfactants, suspending agents, alkaline builders, and optical brighteners. Soaps and detergents have been reported to create both false-negative and false-positive results on several different immunoassays.
Conclusion

The current study concludes that some adulterants make it easy to produce false negative results and the specimen integrity testing is inadequate in detection of these adulterants.

References


تأثير خمسة من المواد الشائبة على الكشف عن الترامادول وقياسه كمية في عينات البول

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المملوكة العربي

المقدمة: تزايدت في الآونة الأخيرة نسب تعاطي الترامادول في بعض دول أفريقيا وغرب آسيا مع الأخذ في الاعتبار ضبط كميات كبيرة من الترامادول في شمال وغرب أفريقيا خاصة في مصر. أصبح اختبار البول لتعاطي المخدرات سلاحًا لا يتجزأ في الحرب ضد المخدرات. ومن المشكلات الكبيرة في جميع اختبارات المخدرات في البول إمكانية غش العينة أو استبدالها للحصول على نتائج سلبية.

الهدف من الدراسة: دراسة التأثير الكمي والنوعي لخمسة مواد شائبة على عينات البول الائتلافية للترامادول.

طريقة البحث: أجريت الدراسة على عينات بول جمعت من المتضررين على معمل السموم الأكليبيكية بجامعة سوهاج. تم اختبار العينات للتآكد من سلامتها عن طريق فحص الام الكهرباوجي والكثافة النوعية والكرياتين. تم اختبار العينات بواسطة جهاز الدناةة لتحديد كمية الترامادول. ثم تم تأكيدها وتحديد كمية الترامادول بواسطة جهاز الكروماتوغرافيا ذات الكفاءة العالية.

النتائج: أظهرت عينات البول المعشوشة بالخل والصابون اليدوي السائل ومنظف الدرانو نتائج سلبية خاطئة عن طريق اختبار الدناةة لتدقيق كمية الترامادول. كما أظهرت جهاز الكروماتوغرافيا ذات الكفاءة العالية في عينات البول المعشوقة بنسبة 0.4% من الخل و0.4% من منظف الدرانو.

الخلاصة: إضافة بعض المواد الشائبة إلى عينات البول الائتلافية للترامادول يؤدي إلى نتائج سلبية خاطئة. وتحذيرات سلامة العينات قد تكون غير كافية للتعرف على هذه المواد.

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