Commercial herbal medicines used as African traditional medicines: Ngoma Herbal Tonic Immune Booster interferes with a rapid urine drug screening test

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Background. The prevalent use of African traditional medicine by the general public has been reported. With commercialisation and marketing, some of the herbal medicines (HMs) used are readily available over the counter, most of them promoted as immune boosters. These commercial HMs have not been taken through clinical trials and other tests that would validate their composition and safety, and other properties such as their effect on laboratory diagnostic tests.

Objective. To investigate the cross-reactivity of selected HMs with commonly tested drugs of abuse (DoA) using a qualitative rapid urinary assay.

Methods. The six HMs selected were bought from local pharmacies. A rapid urinary screening test was performed with the Instant View Multi-Drug of Abuse Test kit from Labstix Diagnostics. Drug-free urine (DFU) was pooled from samples donated by healthy volunteers. Urine samples that had tested positive for DoA were obtained from a pharmacology laboratory. Aliquots of the urine samples were spiked with the HMs in neat and diluted form, and tested at various time intervals.

Results. The results for the DFU samples spiked with the HMs remained negative. There were no significant changes in pH or specific gravity of the samples. The results of samples that had tested positive for tetrahydrocannabinol (THC) were not altered by five of the HMs when spiked at 40% v/v. The HM Ngoma Herbal Tonic Immune Booster caused false-negative results for the THC test.

Conclusion. An important finding is that the herbal mixture Ngoma Herbal Tonic Immune Booster caused false-negative results for the cannabinoid screening test. It adds to the list of substances that may be potential adulterants of urine for screening tests.


The prevalent use of African traditional medicine (ATM) by the general public in South Africa (SA) has been reported.[1] It has been documented that most people consult a traditional health practitioner before a primary health practitioner,[2] and it has been reported that most people consult a traditional health practitioner before a primary health practitioner,[2] and it has been documented that most people consult a traditional health practitioner before a primary health practitioner,[2] and it has been documented that most people consult a traditional health practitioner before a primary health practitioner,[2] and it has been documented that most people consult a traditional health practitioner before a primary health practitioner,[2] and it has been documented that most people consult a traditional health practitioner before a primary health practitioner.[2] As a result of commercialisation and marketing, some herbal medicines (HMs) are readily available over the counter, most of them being promoted as immune boosters. The use of these herbal medicines (HMs) has not been taken through clinical trials and other tests that would validate their composition and safety, and other properties such as their effect on laboratory diagnostic tests.

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There could be more issues concerning the use of HMs by sportspeople. The HMs may have properties similar to some banned substances, so that when tested, a sportsperson could unknowingly test positive for banned substances. Common substances known to cause false-positive results on urine screening tests for drugs of abuse (DoA) include the Vicks inhaler testing positive for amphetamine, non-steroidal anti-inflammatory drugs such as ibuprofen testing positive for barbiturates, and cannabinoids and some fluoroquinolones testing positive for opiates.[6,7] There are no listed compounds related to ATM.

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Another area of concern is adulteration of samples. Adulteration is a process of deliberate interference with the process of specimen collection, transport or analysis, with the intention of avoiding a positive test outcome. In vivo adulteration is the intake of interfering substances such as drugs, or lots of water, before sample collection. In vitro adulteration is when substances are added into the urine sample after collection to alter the results, mainly to cause a false negative. Substances commonly used for adulteration include drug-free human, animal or synthetic urine or assay interferants.[11,12]
Generally, urine is tested routinely for DoA in health facilities to aid the diagnosis of potential abusers and some psychiatry patients on admission. The other important testing is for banned substances in special populations such as those whose work environment requires that they be drug free, and people involved in professional sport. Qualitative screening tests are initially used to test for the presence of the DoA, as they are quick, sensitive, relatively economical and simple to use. Results of screening assays are used to indicate that they be drug free, and people involved in professional sport.

A rapid urinalysis assay based on an immunoassay method is the screening test commonly used for DoA. It is based on the principle of antigen-antibody reaction in which the drug or drug metabolite present in urine will interact with a labelled polyclonal or monocolonal antibody provided. Various substances may interfere with immunoassays by cross-reacting with the test reagents and resulting in false-negative, false-positive or invalid results.

Objective

Literature searches provided no results for cross-reactivity of ATMs, or the influence of ATMs on laboratory tests. This study therefore sought to establish whether any of the commonly used CHMs may affect the outcomes of a rapid qualitative screening test for DoA. It was intended to show whether the presence of the CHMs would alter the results of a rapid urinalysis assay.

Methods

Permission for the study (ref. no. MREC/M/09/2011:PG) was granted by the Sefako Makgatho Health Sciences University (SMU) Research Committee and the Superintendent of Dr George Mukhari Academic Hospital.

The rapid urinalysis assay

The test kit used was the Instant View Multi-Drug of Abuse Test kit from Labstix Diagnostics (SA). It consists of test cassettes, each with six panels for concurrent testing of amphetamine, cocaine, methamphetamine, morphine, tetra-hydrocannabinol (THC) and methylenedioxymethamphetamine. The test is a one-step lateral flow chromatographic immunoassay. Each panel contains drug-protein conjugate immobilised on a porous membrane support. The test is based on the principle of competition for limited binding sites on the antibody between the drug or drug metabolite present in the urine sample and the immobilised drug-protein conjugate. If the drug is absent in the urine, or is present in below cut-off levels, a coloured conjugate will interact with the drug antigen immobilised in the test line and form a coloured line. If the drug or metabolite is present, it will bind to and saturate all the binding sites on the antibody conjugate, preventing any binding by the drug antigen at the test area. The results are negative if two colour lines appear, one in the test area and the other in the control area, regardless of the intensity of the line. The results are positive when only one colour line appears in the control site and are invalid if no colour line appears on the test as well as the control site.

A 10-test dipstick kit from Labstix Diagnostics was used to test qualitatively for pH and specific gravity (SG).

The herbal mixtures and preparation

The six CHMs selected for the study are used as immune boosters or to strengthen the body and are commonly sold in many retail outlets; they were bought from local pharmacies. They were Intlamba Zifo (denoted HM1 for the purposes of this study), Maphilisa HM (HM2), Matla African Medicine for All Diseases (HM3), Ngoma Herbal Tonic Immune Booster (HM4), Stametta Body Healing Liquid (HM5) and Vuka Uphile Immune Booster (HM6). From each of the six CHMs, three serial dilutions of 10, 100 and 1 000 times were made with distilled water. These HMs were taken through the test procedure outlined in Fig. 2.

Testing drug-free urine (DFU)

Ten urine samples were collected from healthy volunteers, each of whom confirmed that they had not taken any drugs, HMs or over-the-counter medicines within the preceding 7 days. The samples were pooled as DFU. Both the individual samples and the DFU were tested with the dipsticks for pH and SG. The remaining pooled DFU was stored in the refrigerator at 4°C.

Four samples of 9 mL each of DFU were made for each of the six CHMs. A volume of 1 mL each of the neat HM and the three dilutions were added to the samples. A dipstick test was done on each spiked DFU sample and the results were recorded. The spiked DFU samples were then tested using the Instant View Multi-Drug of Abuse Test cassette, following the instructions in the test kit manual. The test was repeated at time intervals of 8, 24, 48, 72, 96 and 120 hours and the results were recorded. Following up on the results of the test, six aliquots of the DFU were each spiked with the HMs at 40% v/v (2 400 µL DFU + 1 600 µL CHM). The control for each sample was composed of 2 400 µL DFU plus 1 600 µL of distilled water to make up for the dilution.

RESULTS

Urine testing negative for DoA (DFU)

Spiked with HMs at
- low concentrations (10, 100 and 1 000 × dilutions)
- high concentrations (40% v/v)

Qualitative rapid urine screening test
(Instant View Multi-Drug of Abuse Test)

Urine testing positive for DoA (D-U)

REPORT RESULTS

Fig. 1. Flow diagram for routine urine testing for drugs of abuse. (HPLC = high-performance liquid chromatography; GC-MS = gas chromatography-mass spectrometry).

Fig. 2. Outline of the method used. (DoA = drugs of abuse; DFU = drug-free urine; D-U = positive urine samples; HMs = herbal medicines.)
Testing urine positive for DoA
Urine samples that had been sent to SMU’s Department of Pharmacology and Therapeutics laboratory for routine testing and had tested positive for any of the tested substances were collected. The positive urine samples (D-U) were labelled accordingly, and centrifuged for 5 minutes at 1 000 rpm (250 g). Each supernatant was collected and stored in the refrigerator at 4°C until needed.

Six aliquots of the D-U samples were each spiked with one of the six HMs at 40% v/v. The positive control for each sample was composed of 2 400 µL D-U plus 1 600 µL DFU. These samples were tested following the same procedure according to the test kit instructions.

Results

Testing DFU
The urine samples obtained from healthy volunteers appeared normal. The colour of the samples was various shades of amber, but none of the observed parameters deviated from the normal ranges as indicated by the dipstick test. The pH and SG of the individual urine samples differed, but the pooled sample values were pH 6.0 and SG 1.020.

The results of the rapid qualitative test on the spiked DFU samples were all negative for the tested DoA and remained negative from day 1 till day 5. The dipstick results for pH and SG also remained the same for each of the samples throughout the 5 days of testing.

The results for the 40% v/v spiked urine samples were also negative, and remained negative throughout the 5 days. The mean SG and pH for the DFU spiked with the HMs were 1.020 and 5.5, respectively.

Testing urine that was positive for DoA
Eight urine samples (D-U) that had tested positive for THC were used in this assay. The pH of samples ranged from 5 to 7.5 and the SG from 1.005 to 1.030. The pH of the controls was between 5 and 6.5 and the SG between 1.010 and 1.030. The controls for all the D-U samples remained positive for THC. The results of seven D-U samples spiked with HM1, HM2, HM3, HM5 and HM6 remained positive for THC in all concentrations of the HMs tested, from the lowest to the highest (40% v/v).

The D-U samples spiked with HM4 at the highest concentration (40% v/v) tested negative for THC. Because this HM was the only one that was alcohol based, alcohol had to be excluded as a possible cause of the false-negative result. A 13.5% methanol solution was used to spike some D-U samples, which were then taken through the same procedure. The results of the spiked D-U samples remained positive, confirming that alcohol did not alter the results.

Discussion
Five of the six HMs tested had no influence on the quantitative rapid urinalysis assay. HM4, Ngoma Herbal Tonic Immune Booster, showed false-negative results for THC in the urine samples that were positive for THC when spiked at 40% v/v with HM4. It is stated on the container of this HM that it contains *Sutherlandia*, *Echinacea*, dandelion, alfalfa (lucerne), *Aloe ferox*, *Harpagophytum* (devil’s claw) and alcohol (13.5%). Alcohol was excluded as a possible cause of the false-negative result. The plausible explanation for the false-negative result could therefore be that alcohol as a solvent extracted some lipophilic compounds from the plants used, which would not be extracted by water as a solvent. These compounds would have then interfered with the test reagents, resulting in the false-negative results.

For the test to be positive, the drug metabolites in the urine sample compete for binding on the immobilised protein conjugate (antibody) with the colour-coded drug antigen on the test panel. A negative test result occurs when the colour-coded drug antigen binds to the antibody. The compounds in HM4 interacted with the test reagents and the THC in the sample in such a way that the drug metabolite could not bind to the antibody.

There are several ways in which false-negative results could occur for a particular drug when using immunoassays. A false negative may occur when the drug is present in the sample but the detection limit of the method is too high, or when the actual concentration of the drug in the sample is very low.9,13 Most screening tests have cut-off concentrations above which the results would be reported as positive.14-16 It may also be caused by cross-reactivity of the antibody in the assay, the time lapse between drug ingestion and specimen collection, or adulteration of the sample.17

For HM4, the false-negative results occurred at high concentration (40% v/v). Although the test used is a qualitative test, it has a quantitative part due to the cut-off levels, which give a guide to the possible concentration of the substance if positive.16 The suggestion here is that the presence of HM4 at low levels, which are the most likely in vivo physiological levels, would not affect the screening test. The high concentration used would be most likely to be achieved by *in vitro* sample manipulation, therefore suggesting that HM4 could be an adulterant for THC. Dilution of a sample, which would normally cause reduced concentration to below cut-off levels14 is not a factor in this case because the control sample remained positive.

The false-negative results could also have been caused by cross-reactivity of the antibody in the assay with one or more plant compounds present in the HM. As is well known in HMs, there could be undeclared ingredients in the mixture owing to either deliberate or unintentional contamination,4,17 and also because manufacturers of commercial HMs protect their recipes, as they consider themselves to hold unofficial patents thereof.18 Screening tests are designed to detect classes of drugs, so although they are sensitive they have low specificity.11 THC is a highly lipid-soluble compound of plant origin. It could be that the antibody to THC in the assay has variable degrees of reactivity or affinity towards drugs or metabolites with identical chemical structure to THC, or with unrelated chemical structures. HM4 may contain one or more plant compounds that are structurally related to THC, since THC is of plant origin. Plant compounds that are extracted in alcohol include glycosides, sugars, amino acids, terpenoids, alkaloids and polyphenols.19,20 Whether these were the compounds that interfered with the test is speculative at this stage, as there could be more compounds present. The result cannot be extrapolated to physiological levels, as it was observed at a high concentration (40% v/v) of HM4. However, the result permits the possibility of the risk of the use of HM4 as an adulterant. The cannabinoid assay was recognised to be the assay most susceptible to false-negative results after six out of 16 chemicals commonly used as adulterants reported false-negative results.21 The substances interfering with cannabinoid immunoassays included ibuprofen, tolmetin16 and common household chemicals such as drain cleaner, ammonia and bleach.15,21,22 This is the first report of an ATM, particularly one available over the counter, being a potential adulterant to a cannabinoid urine screening test using a quantitative lateral flow chromatography immunoassay. The outcome also signifies the importance of appropriate labelling of CHMs, which may provide information on the range of possible compounds that may be present in a herbal mixture.

Study limitations
The sample set for this study was small, so a wider study with more samples and replicates and including more herbal medicinal products
should be done. The study method used a rapid screening test from only one manufacturer. The results may therefore not be generalisable to the test kits from other manufacturers.

Conclusion
Of the six HMs tested, only one showed the potential of interfering with a rapid, qualitative urine test for DoA. Ngoma Herbal Tonic Immune Booster caused false-negative results for THC. This is the first finding of the possibility of commercial HMs used as ATMs interfering with rapid, qualitative screening tests. The findings therefore facilitate the route for further studies investigating whether the use of these medicines interferes with diagnostic tests in general.

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Author contributions
MEM conducted the research as part of her PhD studies, and drafted the manuscript. EO and CPKV were co-supervisor and supervisor, respectively; they proofread the draft. All authors approved the final manuscript.

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Conflicts of interest
None


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