Comparative effectiveness of chemical biocides and *Acalypha wilkesiana* leaf extract against postharvest fungal deteriogens of sweet orange (*Citrus sinensis*) fruits

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**Abstract**

This study determined the in-vitro and in-vivo activities of Voriconazole (VOR, 1 μg), Nystatin (NYS, 25 μg), Fluconazole (FLC, 100 μg), red force® (RFC) [Copper (I) oxide, 60% + Metalaxyl-M 6% WP], ultimax plus® (UPL) [Metalaxyl 12% WP + Copper (I) oxide 60% WP], and forcelet® (FCE) [Carbendazim 50% WP] and *Acalypha wilkesiana* (Red Acalypha) on fungal deteriogens obtained from deteriorated sweet orange fruits. The 22 fungal deteriogens tested had in-vitro susceptibility of 13.0 ± 2.6–28.7 ± 1.2 mm inhibition zone for VOR (n = 14), 9.3 ± 1.2–28.0 ± 3.0 mm for NYS (n = 21), 7.7 ± 1.5–21.7 ± 1.5 mm for FLC (n = 4), 13.3 ± 1.2–31.7 ± 2.1 mm for RFC (n = 5), 15.3 ± 4.2–30.7 ± 1.2 mm for UPL (n = 5), 15.7 ± 1.2–31.7 ± 1.5 mm for FCE (n = 19) and 0.0 ± 0.0 mm for Red Acalypha (n = 22). Fifteen of the fungal deteriogens including the yeast, *Candida glabrata* were able to initiate deterioration in-vivo with 40–100% severity within the incubation period of 11 days. FCE biocide significantly (p = 0.001) prevented or reduced deterioration in-vivo. There was no deterioration sign in uninoculated control (Control 1) and uninoculated control with FCE (Control 2). Further search for active agents against fungal deteriogens especially from natural sources is required for longer preservation of *Citrus sinensis*.

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1. Introduction

Citrus is one of the world’s fruit crops of high economic importance with annual worldwide production estimated at over 123 million metric tons as at 2010 [1]. They are the highest valued fruit crop regarding international trade [2]. The commonly cultivated species include sweet orange (*Citrus sinensis*), grapefruit (*Citrus paradise*), lime (*Citrus aurantifolia*), lemon (*Citrus limon*), tangerine (*Citrus reticulate*) and sour orange (*Citrus aurantium*). Oranges constitute the bulk of citrus fruit production, accounting for approximately 55% of global citrus production.

Although believed to have originated from certain parts of Southeast Asia [3,4], tropical Africa is greatly endowed with varied ecological and climatic conditions suitable for the production of citrus in commercial quantity. For instance, Nigeria has an estimated annual citrus production of about 930,000 metric tons [5] which mostly comprised oranges [2,6]. Developing nations, therefore, have the immense comparative advantage and the potential to lead the world in citrus production and trade.

Sweet orange (*Citrus sinensis*) which belongs to the family Rutaceae are globose to ovoid shaped, berry-like edible fruits that vary in their sizes, colours, shapes and fruit quality [7]. Early acceptance of sweet orange in international trade despite their being highly perishable might be due to their high nutritional and therapeutic values [8]. They contain high vitamin C content, appreciable amount of vitamin A, folate, and fibre which have implication in bone formation, eye health, DNA production, reduction in cardiovascular disease risk, etc. They, however, do not contain fat, sodium, cholesterol and has low energy level value [9].

Commercial cultivation of this important fruit crop for large-scale processing and international export is still at infancy in most developing countries. For instance, most of the existing plantations are in mixed cropping system in the remote part of Nigeria’s rainforest and guinea savannah where there is no good transportation system. Only a few large plantations are in existence [2]. According to NIHORT [10], about 45% of citrus currently produced in Nigeria is consumed fresh, 25% is processed while as much as 30% is wasted due to postharvest spoilage mostly caused by fungal...
pathogens. Infection of oranges with fungal species may occur before harvesting, at the time of harvesting and during postharvest handling, storage or distribution [11].

Fungi have been extensively studied as important plant pathogens especially in crops. Several species were implicated in various plant diseases such as Alternaria brown spot, grey spot, melanose, scab disease, black spot, post bloom fruit drop, Pseudocercospora fruit and leaf spot, etc. [12–16]. In the study of Embaby et al. [17], the five species of fungal pathogens, Alternaria citri, Botryodiplodia theobromae, Fusarium sp., Penicillium digitatum and Penicillium italicum isolated from rotten orange fruits were pathogenic and increase considerably the percentage of disease incidence and disease severity as well as significant decrease in shelf life.

Also, fungi associated with spoilage of sweet orange (Citrus sinensis) have been reported to produce toxins [18] and have been reported to be pathogenic [19]. They produce aflatoxins which cause cancer of the liver, aflatoxicosis and acute hepatitis in humans [20]. A single infected orange can be a source of infection to other oranges during storage and in transit [21]. Isolation and identification of the pathogens are desirable to strategize the control measures to (or “intending to”) reducing losses due to spoilage or infections [22].

There has been an increased need to identify and isolate fungi associated with orange spoilage as a proactive measure towards a drastic reduction in economic loss due to fruit decay especially in developing nations where there is no access to improved postharvest storage and transport. The effort may lead to the discovery of previously unknown pathogens that must be isolated and studied.

Previous studies on the fungal postharvest spoilage of sweet orange fruits in Nigeria were limited to isolation, characterization and pathogenicity of the associated fungi [23–28]. Such information may only be considered meagre considering the huge potential for sweet orange production by the citrus producing states in the country. Limited information, however, exists on the susceptibility of the indigenous spoilage-associated fungi to both chemical and biological controls.

Red Acalypha are known to possess antimicrobial activities with extensive usage in the treatment of bacterial and fungal infections [29]. However, its use in prevention of deterioration in sweet orange fruits has not been investigated. Also, there is no report on the comparative evaluation of its effectiveness with chemical biocides in prevention of the activities of fungal deteriogens in sweet orange fruits.

This study, therefore, focused on the characterization of fungi associated with deteriorating sweet orange fruits. In-vitro and in-vivo susceptibility of the fungal deteriogens to chemical biocides and leaf extracts of Red Acalypha were compared with the view to developing a more sustainable biological control for sweet orange deterioration in developing nations.

2. Materials and methods

2.1. Study location and sampling

The study was conducted in the Microbiology Laboratory of Osun State University, Osogbo, Nigeria. Deteriorated sweet orange (Citrus sinensis) fruit samples were obtained from fresh fruit stalls in a local market in Osogbo (7.7667° N, 4.5667° E) during peak citrus fruits production season (August–November 2015). Apparently-healthy ripe, sweet orange fruit samples were obtained from the horticultural garden within the University main campus.

Deteriorated sweet orange fruit samples (10 fruit samples) were randomly collected in sterile polythene bags. As indication of deterioration, sweet orange fruits showing obvious mechanical wound or bruise, with purplish to dark brown rot, blue rot, green rot or black lesions were selectively obtained [30]. Samples were transported immediately to the laboratory and stored refrigerated (4 ± 2°C) until analysed. All apparently-healthy ripe, sweet orange fruit samples (125 fruit samples) were obtained from the same sweet orange tree over a period of two months for the in-vivo assays.

2.2. Isolation and characterization of fungal deteriogens

Deteriorated sweet orange fruit samples were surface-sterilized with cotton wool soaked in 70% ethanol and then rinsed in two different changes of sterile distilled water. Each of the infected sweet orange fruit tissue was cut out into small segments using a sterile scalpel. One gram of the excised portion was aseptically weighed and syringed into a tube of 9 ml of phosphate buffered saline (1X PBS) and shaken vigorously. The tube content then served as stock for 10-fold serial dilution. A 100 µl inoculum each of the 10^-2 and 10^-3 dilutions was aseptically spread-plated onto sterile solidified Potato Dextrose Agar (PDA, Oxoid®) medium containing Chloramphenicol (30 µg/ml) in duplicates.

All the inoculated plates were incubated at 28 ± 2°C for seven days. Fungal isolates were further purified on fresh PDA plates. Pure strains were then maintained on PDA slants and stored in the refrigerator at 4 ± 2°C for further analysis. Purified fungal colonies were identified based on their morphological and microscopic appearances. Growth on PDA plates was observed for appearance and growth pattern, colony texture, colony colour and pigmentation, presence or absence of aerial mycelium, and presence or absence of wrinkles and furrows [31,32].

A drop of lactophenol cotton blue stain was placed on a clean grease-free glass slide after which a sterile inoculating wire loop was used to pick the mycelium from the edge and the centre of each growing fungal colony onto the glass slide [33,34]. The mycelium was then spread evenly on the slide. Teasing was done to separate the mycelium to get a homogenous mixture, and the mixture was then gently covered with coverslips and then allowed to stay for few seconds before observing with the microscope under X40 magnification.

The microscope examination of actively growing mould was by their vegetative and reproductive structures such as hyphal colour and structures, shape and size of conidia, conidiophores, and microsclerotia. Isolates’ macroscopic and microscopic features were then compared with standard keys and published literature [34–38].

2.3. Fungicide preparation

The antifungal susceptibility testing discs of Voriconazole (VOR, 1 µg); Nystatin (NYS, 100 µg); Fluconazole (FLC, 25 µg) were obtained directly from the manufacturer (Oxoid, England). Powdered fungicidal formulations such as red force® (RFC) [Copper(I) oxide, 60% + Metalaxyl-M 6% WP], ultimax plus® (UPL) [Metalaxyl 12% WP + Copper(I) oxide 60% WP], and forcelet® (FCE) [Carbendazim 50% WP] were obtained from local agrochemical stores in Osogbo, Nigeria.

The fresh leaves of Red Acalypha (A. wilkesiana), the extract of which was used in this comparative study, were obtained within the premises of Osun State University, Nigeria. The taxonomy was authenticated by a Plant Biologist in the Department. Extract of this Red Acalypha was prepared in hot water as previously described [39,40].

For comparability, the three powdered fungicidal formulations and the Red Acalypha extract were made into discs to simulate the commercial antifungal susceptibility testing discs. Concentrations
of 1.25 mg, 0.6 mg and 0.6 mg of RFC, UPL and FCE respectively as well as 1% w/v, 0.5% w/v, 0.25% w/v and 0.125% w/v of Red Acalypha leaf extract respectively were prepared. A 100 μl of each solution was impregnated on sterile perforated Whatman filter paper (6 mm) and allowed to dry. All the discs were preserved at 4 °C until used.

2.4. In-vitro susceptibility test of isolated fungal deteriorogens

The disk-diffusion method was used to determine the in-vitro susceptibility of the isolated fungal deteriorogens to the six commercial fungicides (chemical biocides) and the Red Acalypha leaf extract [41]. A 100 μl standardised spore suspension (10⁶ spores/ml) of each isolated fungal deteriorogens was swabbed on the surface of PDA plates. The fungicide discs were aseptically placed on the inoculated plates and incubated at 28 ± 2°C for 48 h. The experiment was done in duplicate, and the test was conducted twice. The zone of inhibition was measured with a scale in mm.

2.5. In-vivo susceptibility test of isolated fungal deteriorogens

Based on the results of in-vitro susceptibility test, only the FCE was used in the subsequent in-vivo susceptibility assay. Apparently healthy sweet orange fruits were used for this assay. The fruits were fresh and slightly ripe (referred to as fresh orange fruits in this study). The fresh orange fruits were surface-sterilized by dipping in 0.4% freshly prepared chlorine for 2 min and subsequently rinsed in three different changes of sterile distilled water [34].

Two different Groups of fresh orange fruits containing two oranges each were used (A and B). Similar experiment was also set up for two different Controls (1 and 2). Two different treatments were administered to each Control and Group experiments as shown in Table 1.

For each Control (1 and 2) and Group (A and B), treatment one was done aseptically by inoculating sterile distilled water or fungal spores of each isolated fungal deteriorogens into two fresh orange fruits using different syringes and needles. For the Control experiments, treatment two was done on each by soaking in sterile distilled water (Control 1) or filter-sterilized FCE (Control 2). However, for the Groups experiments, different containers were used for soaking of oranges for each isolated fungal deteriorogen in sterile distilled water (Group A) or filter-sterilized FCE (Group B).

The treated fresh orange fruits were then placed in clean perforated plastic plates, each moistened with wet balls of absorbent cotton wool to create a humid environment, and incubated at room temperature (28 ± 2°C) for 11 days. Treated samples were left undisturbed for the first four days for visible deterioration signs. Measurement of the diameter of rot was done at days 5, 6, 7, 8, 10 and 11. The percentage of deterioration was then calculated as the percentage of the diameter of the external rotted portion to the total diameter of the sweet orange fruit according to the following formula: % deterioration = r/ N × 100 where r = diameter of the external rotted portion and N = total diameter of the orange [42].

Visual observation of the treated fresh orange fruits was also made each day for deterioration signs and severity of deterioration was based on the extent of spoilage observed on the 11th day of incubation and scored according to the following scales: 0 = no spoilage sign; 1 = spongy appearance or rot; 2 = spongy appearance + rot; 3 = spongy appearance + rot + watery exudate; 4 = spongy appearance + rot + mycelial growth and 5 = spongy appearance + rot + watery exudate + mycelial growth. The percentage deterioration severity was calculated by adapting and modifying the formula of Townsend and Heuberger [43] as follows:

\[
D.S.\% = \frac{\sum (n \times r1) - ... - (n \times r5)}{5N} \times 100
\]

where D.S. = disease severity; n = number of sweet orange fruits used; r1 ... r5 = category number and N = total examined fruits multiplied by the maximum numerical disease grade, i.e. 5. The causal agents (i.e. deteriorogens) were re-isolated from the infected orange fruits and compared with the original fungal isolates at the end of the experiment.

2.6. Statistical analysis

The results of the study are presented with descriptive statistics [mean ± SD]. Inferential statistics to establish the presence or absence of significant difference in mean values were done using ANOVA. A p value of <0.05 was used as indicator of significant difference in a two-tail hypothesis. The analyses were done using SPSS 17.0 software (USA).

3. Results

3.1. Fungal isolates from deteriorated sweet orange fruits

The presumptive identity of the twenty-two (22) fungal species isolated from the deteriorated sweet orange fruits is presented in Fig. 1. They include 20 mould and two yeast groups belonging to 9 different genera namely: Alternaria (1), Arthroderma (1), Aspergillus (6), Candida (1), Fusarium (5), Gliocladium (1), Paecilomyces (1), Penicillium (5) and Trichosporon (1).

Aspergillus niger had the highest occurrence of 15.1% in the deteriorated sweet orange fruits followed by Alternaria sp. and Aspergillus fumigatus with 9.1%, Fusarium merismoides, Fusarium redolens, Gliocladium corda and Penicillium citrinum, occurred in similar proportion of about 6.1% in the deteriorated sweet orange fruits. Arthroderma lenticaliare, Aspergillus alutaceus, A. candidus, A. flavus, A. wentii, Fusarium chlamydosporum, F. sporotrichioides, F. flocciferum, Paecilomyces marquandii, Penicillium griseofulvin, P. canescens, P. chrysogenum, P. frequentans as well as the yeasts Candida glabrata and Trichosporon asahii had the least occurrence 3.0% respectively (Fig. 1).

3.2. In-vitro response of fungal deteriorogens to chemical biocides and Red Acalypha extract

Results of the in-vitro exposure of fungal isolates from deteriorated sweet orange fruits to the six commercial fungicides (chemical biocides) and the Red Acalypha leaf extract are presented in Table 2 while their appearance on plates are also presented in Plate 1.

In-vitro susceptibility of fungal deteriorogens from deteriorated sweet orange fruits to VOR ranged between 13.0 ± 2.6 mm and

| Table 1 | Experimental design for the In-vivo susceptibility test. |
|---------|--------------------------------------------------------|
| Treatment one (by injection) | Treatment two (by soaking for 30 min) |
| Control 1 | Fresh orange fruits + 100 μl sterile distilled water | Sterile distilled water |
| Control 2 | Fresh orange fruits + 100 μl sterile distilled water | Filter-sterilized FCE (1.25 g/l) |
| Group A | Fresh orange fruits + 100 μl fungal spores (0.5 McFarland standard at 520 nm) | Sterile distilled water |
| Group B | Fresh orange fruits + 100 μl fungal spores (0.5 McFarland standard at 520 nm) | Filter-sterilized FCE (1.25 g/l) |
The largest zone of inhibition by VOR was obtained in *Penicillium griseofulvum* (28.7 ± 1.2 mm) and the least in *Penicillium canescens* (13.0 ± 2.6 mm). No zone of inhibition (0.0 ± 0.0 mm) was observed for *Fusarium chlamydosporum*, *Fusarium flocciferum*, *Fusarium merismoides*, *Gliocladium corda*, *Paecilomyces marquandii*, *Penicillium chrysogenum*, *Penicillium citrinum* and *Penicillium frequentans*. VOR was active against 63.6% fungal deteriogens (*n* = 14) including the two yeast species. It was active against 100% of the *Aspergillus* sp. (*n* = 6), 40% of the *Penicillium* sp. (*n* = 5), 40% of the *Fusarium* sp. (*n* = 5), and the *Alternaria* sp. (*n* = 1).

NYS was the most effective antifungal agent inhibiting 21 (95.5%) out of the 22 fungal deteriogens. It was active against 100% of the yeasts species (*n* = 2) and 95% of the mould species (*n* = 20) except *Paecilomyces marquandii*. NYS showed the highest activity against *Penicillium chrysogenum* (28.0 ± 3.0 mm) and the least activity against *Fusarium sporotrichioides* (9.3 ± 1.2 mm).

FLC was the least effective among the commercially available fungicide. It was active against only four out of the 22 fungal deteriogens tested (18.18%). The sensitive species included the moulds *Alternaria* sp. and *Fusarium sporotrichioides* as well as the yeasts *Candida glabrata* and *Trichosporon asahii*. FLC had the highest

![Fig. 1. Presumptive identity and % occurrence of fungal species isolated from deteriorated sweet orange fruits.](image)

| Fungal deteriogens                  | Zone of inhibition diameter [X ± SD], mm |
|------------------------------------|----------------------------------------|
|                                    | VOR         | NYS         | FLC         | RFC         | UPL         | FCE         | Red Acalypha |
| *Alternaria* sp.                   | 23.7 ± 1.2  | 27.7 ± 1.5  | 21.7 ± 1.5  | 0.0 ± 0.0   | 0.0 ± 0.0   | 0.0 ± 0.0   | 0.0 ± 0.0    |
| Arthroderma lenticulare            | 17.0 ± 1.0  | 10.7 ± 0.6  | 0.0 ± 0.0   | 0.0 ± 0.0   | 0.0 ± 0.0   | 15.7 ± 1.2  | 0.0 ± 0.0    |
| *Aspergillus* alutaceus            | 17.7 ± 1.2  | 14.3 ± 1.2  | 0.0 ± 0.0   | 0.0 ± 0.0   | 0.0 ± 0.0   | 31.7 ± 1.5  | 0.0 ± 0.0    |
| *Aspergillus* candidus             | 17.0 ± 2.6  | 13.7 ± 0.6  | 0.0 ± 0.0   | 0.0 ± 0.0   | 0.0 ± 0.0   | 29.3 ± 1.5  | 0.0 ± 0.0    |
| *Aspergillus* flavus               | 18.0 ± 2.0  | 14.3 ± 0.6  | 0.0 ± 0.0   | 0.0 ± 0.0   | 0.0 ± 0.0   | 25.3 ± 1.2  | 0.0 ± 0.0    |
| *Aspergillus* fumigatus            | 17.3 ± 3.1  | 22.3 ± 5.0  | 0.0 ± 0.0   | 0.0 ± 0.0   | 0.0 ± 0.0   | 30.3 ± 1.2  | 0.0 ± 0.0    |
| *Aspergillus* niger                | 25.0 ± 1.7  | 21.3 ± 3.1  | 0.0 ± 0.0   | 0.0 ± 0.0   | 0.0 ± 0.0   | 22.3 ± 2.5  | 0.0 ± 0.0    |
| *Aspergillus* wentii               | 22.3 ± 0.6  | 16.3 ± 1.2  | 0.0 ± 0.0   | 13.3 ± 1.2  | 0.0 ± 0.0   | 26.7 ± 1.2  | 0.0 ± 0.0    |
| *Candida* glabrata                 | 16.7 ± 1.2  | 10.7 ± 1.2  | 20.7 ± 2.3  | 0.0 ± 0.0   | 0.0 ± 0.0   | 26.7 ± 1.2  | 0.0 ± 0.0    |
| *Fusarium* chlamydosporum          | 0.0 ± 0.0   | 17.3 ± 1.2  | 0.0 ± 0.0   | 0.0 ± 0.0   | 0.0 ± 0.0   | 17.3 ± 1.2  | 0.0 ± 0.0    |
| *Fusarium* flocciferum             | 0.0 ± 0.0   | 10.7 ± 0.6  | 0.0 ± 0.0   | 12.7 ± 0.6  | 25.7 ± 0.6  | 0.0 ± 0.0   | 0.0 ± 0.0    |
| *Fusarium* merismoides             | 0.0 ± 0.0   | 16.3 ± 1.2  | 0.0 ± 0.0   | 0.0 ± 0.0   | 20.3 ± 2.0  | 17.3 ± 2.0  | 0.0 ± 0.0    |
| *Fusarium* redolens                | 13.7 ± 1.5  | 17.3 ± 0.6  | 0.0 ± 0.0   | 0.0 ± 0.0   | 0.0 ± 0.0   | 26.7 ± 1.2  | 0.0 ± 0.0    |
| *Fusarium* sporotrichioides        | 19.7 ± 0.6  | 9.3 ± 1.2   | 7.7 ± 1.5   | 0.0 ± 0.0   | 0.0 ± 0.0   | 16.3 ± 1.2  | 0.0 ± 0.0    |
| *Gliocladium* corda                | 0.0 ± 0.0   | 21.7 ± 1.2  | 0.0 ± 0.0   | 0.0 ± 0.0   | 0.0 ± 0.0   | 30.3 ± 0.6  | 0.0 ± 0.0    |
| *Paecilomyces* marquandii          | 0.0 ± 0.0   | 0.0 ± 0.0   | 0.0 ± 0.0   | 0.0 ± 0.0   | 0.0 ± 0.0   | 20.0 ± 2.0  | 0.0 ± 0.0    |
| *Penicillium* canescens            | 13.0 ± 2.6  | 19.3 ± 1.2  | 0.0 ± 0.0   | 19.3 ± 1.2  | 18.7 ± 1.2  | 21.7 ± 0.6  | 0.0 ± 0.0    |
| *Penicillium* chrysogenum          | 0.0 ± 0.0   | 28.0 ± 3.0  | 0.0 ± 0.0   | 31.7 ± 2.1  | 30.7 ± 1.2  | 31.0 ± 1.7  | 0.0 ± 0.0    |
| *Penicillium* citrinum             | 0.0 ± 0.0   | 18.3 ± 0.6  | 0.0 ± 0.0   | 16.0 ± 3.0  | 15.7 ± 2.1  | 29.7 ± 1.5  | 0.0 ± 0.0    |
| *Penicillium* frequentans          | 0.0 ± 0.0   | 21.7 ± 0.6  | 0.0 ± 0.0   | 0.0 ± 0.0   | 15.3 ± 4.2  | 23.3 ± 1.5  | 0.0 ± 0.0    |
| *Penicillium* griseofulvum         | 28.7 ± 1.2  | 20.7 ± 0.6  | 0.0 ± 0.0   | 17.3 ± 0.6  | 0.0 ± 0.0   | 20.7 ± 0.6  | 0.0 ± 0.0    |
| *Trichosporon* asahii              | 18.3 ± 0.6  | 20.7 ± 1.2  | 20.7 ± 0.6  | 0.0 ± 0.0   | 0.0 ± 0.0   | 0.0 ± 0.0   | 0.0 ± 0.0    |

Notes: VOR – Voriconazole, NYS – Nystatin, FLC – Fluconazole, RFC – red force®, UPL – ultimax plus®, FCE – forcelet®. [Data represents mean of triplicate analysis].
activity against *Alternaria* sp. (21.7 ± 1.5 mm) and the lowest activity against *Fusarium sporotrichioides* (7.7 ± 1.5 mm).

Only five out of the 22 fungal deteriorogens (22.73%) isolated from the deteriorated sweet orange fruits were sensitive to RFC and UPL powdered biocides. The yeast species, *Candida glabrata*, and *Trichosporon asahii* showed resistance to the two fungal biocides. Up to 80% of the *Penicillium* sp. (*n* = 5) and 16.67% of the *Aspergillus* sp. (*n* = 6) were susceptible to RFC with *Penicillium chrysogenum* and *Aspergillus wentii* showing the highest and lowest sensitivity of 31.7 ± 2.1 mm and 13.3 ± 1.2 mm respectively. Conversely, UPL activity was observed against 80% of the *Penicillium* sp. (*n* = 5) and 20% of the *Fusarium* sp. (*n* = 5) with *Penicillium chrysogenum* and *Penicillium frequentans* showing the largest (30.7 ± 1.2 mm) and the lowest (15.3 ± 4.2 mm) zones of inhibition respectively.

Out of the 22 fungal deteriorogens of the sweet orange fruits tested, 19 which represent about 86.4% were susceptible to the FCE, making it the most effect powdered fungal biocide with field application potential. The FCE was active against 100% of the *Arthroderma* sp. (*n* = 1), *Aspergillus* sp. (*n* = 6), *Fusarium* sp. (*n* = 5), *Gliocladium* sp. (*n* = 1), *Paecilomyces* sp. (*n* = 1) and *Penicillium* sp. (*n* = 5). However, *Alternaria* sp. and the two yeast isolates, *Candida*...
glioblastoma and Trichosporon asahii, were resistant to FCE. Aspergillus alutaceus (31.7 ± 1.5 mm) had the highest zone of inhibition while, the least zone of inhibition was observed in Arthroderma lenticulare (15.7 ± 1.2 mm).

The zone of inhibition for all the fungal deteriogens exposed to various concentrations of Red Acalypha extract was 0.0 ± 0.0 mm. The result showed that Red Acalypha did not show or possess any bioactivities against moulds and yeasts.

3.3. In-vivo pathogenicity of fungi isolated from deteriorated sweet orange fruits

Ability of the 22 fungal deteriogens to initiate visible deterioration in fresh orange fruits and uninoculated control was observed over a period of 11 days in a pathogenicity test (Fig. 2).

Fresh orange fruits inoculated with fungal deteriogens that were capable of causing infection started exhibiting visible deterioration signs measurable in % deterioration (mean value) of rot on day five; Aspergillus candidus (0.8%), Aspergillus wentii (13.1%), Aspergillus flavus (0.6%), Penicillium chrysogenum (5.5%) and Gliocladium corda (47.8%) in day 10; and Penicillium chrysogenum (66.9%) in day 11. However, Alternaria sp., Fusarium chlamydosporum, Fusarium flocciferum, Fusarium merisoides, Fusarium redolens, Paecilomyces marquandii, Trichosporon asahii and uninoculated fresh orange fruits (Control 1) did not show any sign of deterioration within the incubation period of 11 days.

The mean % deterioration of inoculated sweet orange fruits ranged from 6.4 ± 0.0% [mean ± SD] for Aspergillus alutaceus to 98.4 ± 0.1% for Aspergillus candidus. Overall comparison of the mean % deterioration i.e. the indicator of fungal pathogenicity induced by the deteriogens showed highly significant difference (p = 0.001, n = 15). Pairwise comparison of the mean % deterioration revealed that Aspergillus candidus and Aspergillus niger induced significantly higher (p = 0.001) deterioration compared to each of the remaining 13 fungal deteriogens. However, there was no significant difference (p = 0.35; 0.82, 0.77 and 0.45 respectively) when in-vivo pathogenicity of Aspergillus wentii was compared with those of Aspergillus alutaceus, Aspergillus fumigatus, Fusarium sporotrichioides and Penicillium frequentans.

3.4. In-vivo susceptibility of fungal deteriogens to the FCE biocide [Carbendazim 50% WP]

The in-vivo response of the isolated fungal deteriogens inoculated into fresh orange fruits to the FCE biocide is presented as % deterioration and severity on day 11 after inoculation and treatment (Table 3). Representative fresh orange fruit samples inoculated with fungal deteriogens and treated with FCE biocide are also presented in Plate 2.

Antifungal effect of FCE biocide revealed wide differences in susceptibility of each deteriogen. The susceptibility of the fungal deteriogens ranged from 100% for Arthroderma lenticulare, Aspergillus fumigatus, Candida glabrata, Fusarium sporotrichioides, Gliocladium corda and Penicillium griseofulvum to 37.3% for Aspergillus niger.

Compared to Group A, the FCE biocide significantly (p = 0.001) prevented deterioration in fresh orange fruits inoculated with Arthroderma lenticulare, Aspergillus fumigatus, Candida glabrata, Fusarium sporotrichioides, Gliocladium corda and Penicillium griseofulvum while the FCE biocide significantly (p = 0.001) reduced deterioration in fresh orange fruits inoculated with Aspergillus alutaceus, Aspergillus candidus, Aspergillus flavus, Aspergillus niger, Aspergillus wentii, Penicillium canescens, Penicillium chrysogenum, Penicillium citrinum and Penicillium frequentans.

Deterioration severity was highest (100%) in Aspergillus candidus, Aspergillus niger, and Penicillium griseofulvum inoculated fresh orange fruits and was reduced to 20% in Aspergillus candidus, and 0% in Penicillium griseofulvum inoculated fresh orange fruits when treated with FCE biocide. However, there was no reduction in deterioration severity of FCE biocide treated fresh orange fruits inoculated with Aspergillus niger.
Deterioration severity of 80% was reported for fresh orange fruits inoculated with *Aspergillus flavus* and *Penicillium canescens* while 40% was reported for *Aspergillus alutaceus* and *Penicillium frequentans*. Deterioration severity was 60% in fresh orange fruits inoculated with *Candida glabrata*, 40% in *Arthroderma lenticulare*, *F. sporotrichioides*, *G. corda* and 0% when the fresh orange fruits were treated with FCE solution. Deterioration severity of fresh orange fruits inoculated with *Penicillium citrinum* (40%) was reduced to 20% when treated with FCE biocide solution.

There was comparable (p = 1.00) absence of deterioration in the fresh orange fruits + *Arthroderma lenticulare* + FCE (Group B), fresh orange fruits + sterile distilled water (Control 1) and fresh orange fruits + FCE (Control 2). Group B showed significant (p = 0.001) reduction in deterioration compared to Group A (Table 3).

### Table 3

| Fungal Deteriogen | % Deterioration Group A | % Deterioration Group B | % Severity Group A | % Severity Group B |
|-------------------|-------------------------|-------------------------|-------------------|-------------------|
| **Control 1**     | 0                       | 0                       | 0                 | 0                 |
| **Control 2**     | 0                       | 0                       | 0                 | 0                 |
| **Alternaria sp.**| 0                       | 0                       | 0                 | 0                 |
| **Arthroderma lenticulare** | 76.1                   | 0                       | 40                | 0                 |
| **Aspergillus alutaceus** | 6.4                    | 3.4                     | 40                | 40                |
| **Aspergillus candidus** | 58.4                   | 5.5                     | 100               | 20                |
| **Aspergillus flavus** | 45.5                   | 3.9                     | 80                | 80                |
| **Aspergillus fumigatus** | 8.9                    | 0                       | 40                | 0                 |
| **Aspergillus niger** | 89.2                   | 62.7                    | 100               | 100               |
| **Aspergillus wentii** | 8.4                    | 7.2                     | 40                | 40                |
| **Candida glabrata** | 45.3                   | 0                       | 60                | 0                 |
| **Fusarium chlamydosporum** | 0                      | 0                       | 0                 | 0                 |
| **Fusarium flocciferum** | 0                      | 0                       | 0                 | 0                 |
| **Fusarium merismoides** | 0                      | 0                       | 0                 | 0                 |
| **Fusarium redolens** | 0                      | 0                       | 0                 | 0                 |
| **Fusarium sporotrichioides** | 9.1                    | 0                       | 40                | 0                 |
| **Gliocladium corda** | 61.3                   | 0                       | 40                | 0                 |
| **Paecilomyces marquandii** | 0                      | 0                       | 0                 | 0                 |
| **Penicillium canescens** | 36.4                   | 5.1                     | 80                | 80                |
| **Penicillium chrysogenum** | 66.9                   | 6.9                     | 0                 | 0                 |
| **Penicillium citrinum** | 42.2                   | 6.6                     | 40                | 20                |
| **Penicillium frequentans** | 6.9                    | 5.5                     | 40                | 40                |
| **Penicillium griseofulvum** | 33.5                   | 0                       | 100               | 0                 |
| **Trichosporon asahii** | 0                      | 0                       | 0                 | 0                 |

Notes: Group A – fresh orange fruits inoculated with fungal spores only; Group B – fresh orange fruits inoculated with fungal spores and treated with FCE; Control 1 – fresh orange fruits inoculated with sterile water; Control 2 – fresh orange fruits inoculated with sterile water and treated with FCE; *Data represents average values of duplicate samples.

4. Discussion

Postharvest loss of sweet orange (Citrus sinensis) fruits due to the activities of deteriogens can be up to 25% of a single harvest in the developing countries [42]. In developing countries, percentage loss may increase as a direct consequence of the handling, transport and storage practices which increase the physical damage and introduce more microbial deteriogens into the orange fruits. The intrinsic properties of the sweet orange fruits including low pH as well as high sugar and nutrient contents make them more susceptible to fungal attack.

The 22 fungal deteriogens isolated from deteriorated orange fruit samples belonged to the nine genera *Alternaria*, *Arthroderma*, *Aspergillus*, *Candida*, *Fusarium*, *Gliocladium*, *Paecilomyces*, *Penicillium*, and *Trichosporon*. The fungal genera have been found to be associated with spoilage of fruits as previously reported [23,24,27,42,44–46]. Fungal species isolated from this study have been implicated in fruit infection commonly referred to as blue mould, green mould, brown spot, black spot, black rot, black pit, etc. [26].

Aspergillus sp, Penicillium sp. and *Fusarium* sp. which were prevalent in this study are known to be the most important deteriogens affecting fruits worldwide [46]. The high prevalence of *Aspergillus niger* in the fungal isolates obtained in this study is similar to the works of Akintobi et al. [25] and Onuorah et al. [28]. Khan et al. [46] stated that *Aspergillus niger* is the most predominant fungal isolate associated with the deterioration of citrus fruit.

Voriconazole (VOR), Nystatin (NYS) and Fluconazole (FLC) are chemical biocides (synthetic antifungal agents) commonly used for treatment of fungal infections. The in-vitro exposure of the 22 fungal deteriogens to these antifungal agents revealed their broad spectrum nature and therefore confirmed their efficacy as antifungal drugs for treatment of mild to serious invasive fungal infections. More of the fungal deteriogens used in this study were more susceptible to NYS and VOR compared with FLC. VOR is one of the new generation triazoles reported to have a wide range of activity, since they are active against *Candida, Aspergillus, Fusarium, Penicillium, Scedosporium, Acremonium*, and *Trichosporon*, and dimorphic fungi, dermatophytes, and *Cryptococcus neoformans* [47–49].

Red force® (RFC), ultimax plus® (UPL) and forcelet® (FCE) are also chemical biocide brands with antifungal activities usually in the form of powdered fungicide formulations. In-vitro exposure of the 22 fungal deteriogens also revealed that the three chemical powdered formulations have a wide range of antifungal activities with FCE as the most effective. FCE contains Carbendazim, a systemic benzimidazole, commonly used to control a broad range of fungal attacks in crops, fruits and vegetables as well as in their postharvest storage [50]. It has a widespread usage in postharvest preservation of sweet orange fruits, especially in developed countries.

Antimicrobial activities of crude leaf extracts of Acalypha wilkesiana (Red Acalypha) is well documented [51–53]. For instance, Alade and Irobi [51] showed that the water and ethanol extracts of Red Acalypha leaf inhibited the growth of standard and local strains of bacteria and fungi including the yeast *Candida albicans*.
and mould *Aspergillus flavus*. The present study, however, showed that all the 22 fungal deteriogens tested including the yeasts *Candida glabrata* and *Trichosporon asahii* were not inhibited *in-vitro* by the crude extract of Red Acalypha leaf. It, therefore, suggests that extracts of Red Acalypha may not be a suitable biocontrol alternative for prevention of fungal deteriogens in postharvest storage of sweet orange fruits.

The *in-vivo* pathogenicity data (Fig. 2) suggests that a break in the lines of defence of the sweet orange fruits (i.e. skin, flavedo, and albedo) precedes early appearance of deterioration signs and eventual spoilage. Nune et al. [54] stated that wound infection in crop produce is essential to the establishment of deterioration. Production of toxins and important digestive enzymes by the growing fungal deteriogens in the sweet orange fruits might be responsible

**Plate 2.** Representative fresh orange fruits showing deterioration signs after 11 days of incubation. Group A – fresh orange fruits inoculated with fungal spores only; Group B – fresh orange fruits inoculated with fungal spores and treated with FCE; 1 – Control 1 (fresh orange fruits inoculated with sterile water); 2 – Control 2 (fresh orange fruits inoculated with sterile water and treated with FCE); 3 – *Aspergillus flavus*; 4 – *Aspergillus niger*; 5 – *Candida glabrata*; 6 – *Penicillium chrysogenum*. 
for the observed deterioration by 15 of the 22 study fungal deteriorogens [26]. The result also showed that not all the fungal isolates were capable of initiating deterioration in healthy fresh orange fruits. The possible reasons for their non-pathogenicity is subject to further studies.

Deterioration signs were not noticed in the fresh orange fruits inoculated with spores of the fungal deteriorogens until five days of incubation, and where visible deterioration occurred, the rot diameter increased daily until the experiment was terminated on day 11. The reason for the five-day incubation period for the visible development of deterioration was that adequate time is required for the germination of conidia as well as the elongation of fungal hyphae especially of slowly growing fungi [55]. The uninoculated fresh orange fruits used as control in the pathogenicity test remained wholesome throughout the incubation period.

Aspergillus niger was the most pathogenic among the 22 fungal isolates. Similar observation was made by Akintobi et al., [25]. FCE biocide was capable of controlling the development of deterioration caused by fungal isolates in almost all the citrus fruit studied. Abano and Sam-Amoah [56] stated that dips of biocides were effective in controlling post-harvest rot caused by fungal species.

The in-vivo experiment involving FCE biocide treatment of the fresh orange fruits inoculated with various fungal deteriorogens confirmed the broad spectrum nature of the powdered biocide (Table 3). The FCE significantly prevented or reduced deterioration in healthy fresh orange fruits. Use of FCE biocide for preservation of sweet orange fruits is therefore recommended to extend shelf life of sweet oranges. It will thereby reduce postharvest loss of this important seasonal and perishable fruit.

However, no deterioration signs were observed in fresh orange fruits + sterile distilled water (control 1) and fresh orange fruits + FCE (control 2) within the incubation period of 11 days. Absence of deterioration in the two controls confirmed the need for good agricultural practices that will reduce physical damage and contamination of fresh orange fruits, especially during harvest, transport and storage.

5. Conclusion

Sweet orange (Citrus sinensis) fruits is a perishable agricultural produce which is available in large quantity especially during its season. Preservation for a longer period is a major challenge especially in developing countries where farmers do not have access to high-tech harvesting and storage facilities. This study compared the effectiveness of selected chemical biocides and Acalypha wilkesiana leaf extract against postharvest fungal deteriorogens of sweet orange (Citrus sinensis) fruits for sustainable application.

Fungal species are associated with deterioration of sweet orange fruits, and physical damage is a precondition for initiation of biological decay. Although VOR antibiotic possessed highly effective activity against most of the fungal deteriorogens of sweet orange fruits, their use in sweet orange fruits preservation is however limited to avoid early fungal resistance. Forcelet (FCE) powdered biocide, a brand of Carbendazim (50% WP), also possessed a broad spectrum activity (in-vitro and in-vivo) against fungal deteriorogens of sweet orange fruits. In an attempt to provide a cost-effective and sustainable preservation option, Acalypha wilkesiana (Red Acalypha), a suggested biocontrol alternative, did not show any activity against fungal deteriorogens including yeasts.

Although FCE biocide can reduce deterioration in healthy fresh orange fruits inoculated with fungal deteriorogens, maintaining the integrity of the sweet orange fruits especially during harvest, transport and storage is important to the prolonged shelf life and reduction of postharvest loss. Future research will be aimed at testing other non-chemical preservation alternatives for a more sustainable solution.

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References

[1] UNCTAD. Market information in the commodities area: information on citrus fruit. United Nations Conference on Trade and Development. Available from: http://wwwunctad.info/en/Infocomm/Agricultural_Products/Citrus-fruit/market/Production/. Accessed 2015 November; 2013.
[2] Olile IC, Ibeagha OA, Osowalu AF. Citrus fruits value chain development in Nigeria. J Biol Agric Health 2015;5(4):36–47.
[3] Gmitter FG, Hu XL. The possible role of Yunnan, China, in the origin of contemporary citrus species (Rutaceae). Econ Bot 1990;44:267–77.
[4] Liu Y, Heying E, Tamashiroho SA. History, global distribution, and nutritional importance of citrus fruits. Comp Rev Food Sci Food Safety 2012;11(6):530–40.
[5] FAO. High-Level Conference on World Food Security: The Challenges of Climates Change and Bioenergy. Italy: FAO; 2008.
[6] Bates RP, Morris JR, Grandel PG. Principles and practices of small- and medium-scale fruit juice processing. FAO Agricultural Services Bulletin 146. Rome, Italy: Food and Agricultural Organization of the United Nations; 2001; pp. 226, ISBN 92-5-104661-1.
[7] Parle M, Chaturedi D. Orange: range of benefits. Int Res J Pharm 2012;3(7):59–63.
[8] Gorinstein S, Martin-Belloso O, Park Y, Harunenkit R, Lokej A, Ciz M, et al. Comparison of some biochemical characteristics of different citrus fruits. Food Chem 2001;74(3):309–15.
[9] Turner T, Burri BJ. Potential nutritional benefits of current citrus consumption. Agriculture 2013;3:170–87.
[10] NIHORT. 25 Years of Research into Horticultural Crops Development in Nigeria (1975–2000). In: Denton OA, Alasiri KO, Adejojo MA, editors. National Horticultural Research Institute; 2000.
[11] Barth M, Hankinson TR, Zhuang H, Breidt F. Microbiological spoilage of fruits and vegetables. In: Sperber WH, Doyle MP, editors. Compendium of the Microbiological Spoilage of Foods and Beverages, Food Microbiology and Food Safety. Springer Science Business Media: LLC; 2009. p. 135–83.
[12] Wellings CR, Pathogenicity of fungi associated with a citrus greasy spot in New South Wales. Trans Br Mycol Soc 1981;76(3):405–9.
[13] Singh V, Deverall BJ. Boculius subtilis as a control agent against fungal pathogens of citrus fruit. Trans Br Mycol Soc 1984;83(3):487–90.
[14] Chalutz E, Wilson CL. Postharvest biocontrol of green and blue mould and sour rot of citrus fruit by Debaryomyces Hansenii. Plant Dis 1990;74(2):134–7.
[15] Brownning HW, McCrovern RJ, Jackson KK, Calvet DV, Wardawski WF. Florida citrus diagnostic guide on citrus and control green and blue moulds of citrus. Biol Control 1995;5:81–8.
[16] El-Gaouthour A, Wilson CL, Winnieh M, Droyb S, Smilanick JL, Koren L. Biological control of postharvest disease of citrus fruits. In: Gnanamanickam SS, editor. Biological control of crop diseases. Mared Dekker Inc: New York; 2002. p. 289–312.
[17] Embaby EM, Hazaa M, Haqag LF, Ibrahim TE, Abd El-Azem FS. The decay of some citrus fruit quality caused by fungi. J Appl Sci Res 2013;9(11):5920–9.
[18] Al-Hindi RR, Al-Najada AR, Mohammed SA. Isolation and identification of some fruit spoilage fungi: screening of plant cell wall degrading enzymes. Afr J Microbiol Res 2011;5(4):443–5.
[19] Monso EM. Occupational asthma in greenhouse workers. Curr Opin Pulm Med 2004;10(2):147–50.
[20] Bayewu RA, Amusa NA, Ayoola OA, Babalola MO. Survey of the postharvest diseases and aflatoxin contamination of marketed pawpaw fruit (Carica papaya L.) in South Western Nigeria. Afr J Agric Res 2007;2(4):178–81.
[21] Jay JM. Microbial spoilage of food. Modern food microbiology. New York: Chapman and Hall Inc.; 2003. p. 181–95.
[22] Singleton L, Mihaly JD, Rutjes EM. Methodology for research on soil-borne phytopathogenic fungi. St Paul, MN, USA: American Phytopathological Society Press; 1992. p. 264.
[23] Saka A, Mukhtar BQ, Aden A. Isolation and identification of postharvest spoilage fungi associated with sweet orange (Citrus sinensis) traded in Kano metropolis. Bayero J Pure Appl Sci 2009;2(1):122–4.
[24] Akinnusire OO. Fungal species associated with the spoilage of some edible fruits in Maiduguri northern eastern Nigeria. Adv Environ Biol 2011;5(1):157–61.
[25] Akintobi AO, Okonkwo JO, Agbunlo MO, Akano OR, Onianwa O. Isolation and identification of fungi associated with the spoilage of some selected fruits in Ibadan, Southwestern Nigeria. Academic Arena 2011;3(11):1–10.
