Antifungal efficacy of *Thevetia peruviana* leaf extract against *Alternaria solani* and characterization of novel inhibitory compounds by Gas Chromatography-Mass Spectrometry analysis

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**ABSTRACT**

*Alternaria solani*, a plant pathogenic fungus causes significant economical losses of potato crop. The disease is controlled primarily through some traditional methods and most commonly via the application of chemical fungicides. Fungicides treatment is not protected as chemicals pollute environment, effect health vulnerability in humans and when these harmful chemicals enter into the food chain become hazardous to all living entities. Recent efforts have focused on developing environmentally safe, long-lasting, and effective biocontrol methods for the management of plant diseases. Present research focus on screening of crude and partially purified leaf extract of *Thevetia peruviana* for the presence of antifungal efficacy against *Alternaria solani*. It was observed that 100% alcoholic crude and alcoholic fraction of partially purified extract showed maximum inhibitory activity which is due to the presence of different secondary metabolites, revealed by phytochemical screening. Active column fraction (possess best antifungal activity against *Alternaria solani*) was subjected to Gas Chromatography-Mass Spectrometry (GC-MS) analysis. On the basis of peaks matching of GC-MS chromatogram with available data base showed the presence of benzoic acid and oxo-benzoate in active fraction of *Thevetia peruviana* leaf extract which is already known chemical among the phytochemicals described for antimicrobial activity. Further research on development of herbal formulation from the same would be very helpful environment friendly approach to manage concern crop disease.

1. Introduction

Plants have capability to produce a great number of various bioactive compounds like essential oils, gums and resins etc. that have been revealed to possess biological activity against plant fungal pathogens *in vitro* and *in vivo* and can be used as bio-fungicidal products [1]. These plants derived bioactive compounds are produced as secondary metabolites and remain accumulated in different plant parts. The presence of such phytochemicals is a marker that the plant can be a prospective store of these precursors in the development of herbal formulation [2]. These phytochemicals are usually supposed to be more satisfactory and less unsafe for the ecosystems and could be used as substitute remedies for treatment of plant diseases [3]. In the recent years, research work on decoction of biologically active compounds from plant parts and their utilization for therapeutic purposes has increased intensively [4]. Screening of plant parts for the presence of antimicrobial activity is usually done by first preparing plant extract using cold and hot extraction methods. Primary evaluation of antifungal activity commonly done with 100% alcohol, 50% alcohol and 100% aqueous extracts. The secondary metabolites which could be present in the extracts can be isolated in various solvents on the basis of their solubility in the solvents used. Various organic solvents are used on the basis of their polarity in hot extraction using soxhlet assembly for successive extraction.

Early blight is one of the most important diseases which are caused by the fungus *Alternaria solani*. Jones and Grout, that occurs in most potato growing regions worldwide [5]. The most common and effective method for the control of early blight is through the application of foliar fungicides, but the fungicides treatment is not protected as chemicals pollute environment, effect health vulnerability in humans and when these harmful chemicals enter into the food chain become hazardous to all living entities.

Natural plant products have a narrow target range with specific mode of action, therefore are suitable for a specific target, mostly nontoxic for antagonistic microorganisms, show limited field
2. Materials and methods

2.1. Crude and partially purified extracts preparations

Fresh leaves of *Thevetia peruviana* were collected from Botanical garden of Mohanlal Sukhadia University, Udaipur Rajasthan, India. The plant was submitted for identification at Herbarium of Department of Botany, University of Rajasthan, Jaipur, India. *Thevetia peruviana* Linn. Vide voucher specimen number RUBL 211651. The collected plant material was shade dried at room temperature. It was then ground in an electrical grinder. The ground material was passed through a sieve of mesh with size 60 mm to obtain a fine powder. It was then used to prepare the extract. Crude extract was prepared according to the cold extraction method. Cold extraction was prepared in water, 50% alcoholic alcohol and absolute alcohol. 20 gm of dried and powdered plant material was suspended in 100 ml water, a mixture of 50 ml water and 50% alcohol and 100 ml absolute alcohol separately, and kept for 48 h. After 48 h each mixture was filtered through Whatman filter paper no.1. And filtered material was vacuum dried using rotary vacuum evaporator (Remi, India). The dried residue was used as extract and solvent was recycled [9].

2.2. Antifungal activity of crude and partially purified fraction of *Thevetia peruviana* (pers.) schum. leaf extract

Poison food technique [10] was used for the examining the antifungal activity of crude and partially purified fractions of *Thevetia peruviana* leaf extract. In this respect 100 mg of extract was dissolved in 10 ml acetone to prepare stock solution of 10 mg/ml concentration. The Petri-plates inoculated with tested fungi containing extract were incubated at 28 ± 2 °C for seven days. Mancozeb, Bavistin, and only PDA culture media were used as control series along with test samples. Antifungal activities of extracts were measured as a function of increasing or decreasing the growth of 6 mm disc of each inoculum. After seven day of incubation the average diameter of the fungal colonies was measured and mycelial growth in percentage was calculated by the following formula:

\[ \text{Mycelium growth inhibition} = \frac{G_c - G_t}{G_c} \times 100 \]

Where, Gc, growth of mycelia of control; Gt, growth of mycelia of treatment.

2.3. Effect of extracts on morphology of test fungi

Effect of extract on a variety of morphological and cytological parameters of test pathogen was studied. Test fungi were treated with rising concentrations from 1.25 mg/ml to 10 mg/ml of the extract up till MIC. A small fungal biomass containing mycelium, vesicle and conidia/spores were separated from each tube and microscopic assessment was made after staining with cotton blue and mounting in lacto-phenol. Alteration in mycelium width, conidia size morphology were recorded with the help of Olympus trinocular research microscope BX-51 and analyzed by Image analysis software Olysys Bioreport 3.2 of Olympus. Conidia counting were done by haemocytometer.

2.4. Phytochemical analysis of leaf extracts of *Thevetia peruviana* Linn

Qualitative methods were used for the identification of different secondary metabolites or phytochemicals present in the plant extracts. Various fractions of leaf obtained by successive extraction were then subjected to qualitative test suggested by Kokate et al. [11]. The extract was analyzed for the presence and absence of alkaloids, steroids, volatile oils, carbohydrates, tannins, flavonoids and saponins.

2.5. Tests for detection of secondary metabolites

Phytochemistry of plant extracts was also studied by rapid qualitative tests [12].

2.6. Alkaloids

Alkaloids are heterocyclic nitrogenous compounds which are found naturally in plants. Structurally have one or more nitrogen which hold heterocyclic ring. Mayer’s test or Wagner’s test or Hager’s test used to find out the occurrence of alkaloids in the partially purified fractions. A cream coloured precipitate is produced in the reaction with Mayor’s reagent; Wagner’s reagent results in formation of reddish brown precipitate whereas Hager’s reagent gives yellow precipitate. After adding few drops of diluted HCL in the little quantity of extract it was stirred and filtered. Various alkaloid reagents were tested on the filtrate and development of coloured precipitate was observed.

2.7. Volatile oils

Volatile or essential oils are the aromatic volatile chemical constituents of plants. Sudan III test can be employed to identify occurrence of
volatile oils. Red colour was observed, on mixing with Sudan III which reveals the presence of volatile oils. Little quantity of extract and Sudan III dye are mixed for the examination of the development of red colour.

2.8. Tannins

Chemically, tannins have a combination of complex organic substances. Polyphenols are found in these substances. The occurrence of condensed tannins is showed by the appearance of green colour and the existence of hydrolysable tannins is shown by blue colour. Small quantity of extract was used and treated with alcohol FeCl₃ solution and observed for colour development.

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Table 3
Antifungal activity of standard fungicides with water control *Alternaria solani*.

| No. | Standard fungicides and water control | Growth Diameter after 7 days (mm) | % Mycelial growth inhibition |
|-----|-------------------------------------|----------------------------------|-----------------------------|
| 1   | Mancozeb                           | 16.00 ± 1.00                     | 80.16                       |
| 2   | Bavistin                            | 33.66 ± 0.57                     | 58.26                       |
| 3   | Water (control)                     | 80.66 ± 0.57                     | No inhibition               |

**Fig. 1.** Antifungal activity of *Thevetia peruviana* against *Alternaria solani*. A. Crude extracts [1] 100% alcoholic [2] 50% alcoholic [3] 100% aqueous, B. Partially purified fractions [1] Petroleum ether fraction [2] Benzene fraction [3] Chloroform fraction [4] Acetone fraction [5] Alcohol fraction [6] Methanol fraction [7] Aqueous fraction. C. Antifungal activity of standard fungicides with water control against *Alternaria solani* [1] Mancozeb [2] Bavistin [3] Control (only PDA media).
2.9. Saponin

Saponins are complex glycoside compounds. In these compounds aglycone is steroidal in nature. Foam test can be employed to identify existence of saponins. Small quantity of extract diluted with 20 ml of distilled water was shaken in graduated cylinder for 15 min. Development of a layer of foam at surface indicates presence of saponins.

2.10. Carbohydrates

Carbohydrates are extensively distributed in plants. Presence of carbohydrate can be detected by Molisch’s test or Fehling’s test. Small quantity of extract was mixed with 5 ml of distilled water and filtered after dissolution. A few drops of naphthol and H₂SO₄ were mixed to the filtrate and the colour of the filtrate changed to purple. It shows the presence of sugar. Likewise, when small amount of filtrate was heated with equal quantity of Fehling A and Fehling B solution, the colour changed to brick red colour. It indicates presence of carbohydrates.

2.11. Flavonoids

Flavonoids are frequently found in plants as glycosides. Structurally, one or more of phenolic hydroxyl groups are combined with sugar residues. Alkaline reagent test is employed to identify flavonoids. Standard test for detection is, mixing of aqueous NaOH with a small quantity of extract. Development of red brown colour shows presence of flavonoids.

2.12. Sterols

Sterols are triterpenes. Structurally sterols are based on cyclopentaneperhydroxyphenanthrene (sterane) ring system. They are also recognized as phytosterols. Liebermann’s Burchard test is used for revealing the presence of phytosterols in extract. Small quantity of extract was mixed with 1 ml of acetic anhydride and 2 ml CHCl₃ followed by gradual addition of concentrated H₂SO₄ through the side of the test tube. Development of a ring of brown colour at junction of two layers is the indication of the presence of sterols.

2.13. Isolation and characterization of active principle from selected leaf extract by using various chromatography techniques

Isolation of active principle from acetone extract of T. peruviana leaf was done by thin layer chromatographic fingerprinting, column chromatography and Gas chromatography. Characterization of the active compound was done by Mass spectrometry analysis of the extracted phytochemicals.
Table 7
RI of TLC finger printing of alcohol fraction of *Thevetia peruviana* leaf extraction.

| Band Number | Colour of Bands | RI value |
|-------------|-----------------|----------|
| 1.          | Blue green      | 0.10     |
| 2.          | Dark blue       | 0.16     |
| 3.          | Yellow          | 0.33     |
| 4.          | Yellow-blue     | 0.44     |
| 5.          | Purple          | 0.56     |
| 6.          | Purple          | 0.70     |
| 7.          | Yellow          | 0.80     |

+ve = Presence, -ve = Absence.

2.14. TLC finger printing of alcohol fraction of *Thevetia peruviana* Linn

For TLC Finger printing of Alcohol Fraction of *Thevetia peruviana* Linn., 10 cm long TLC plate was cut and marked carefully. 10 μl of plant extract was spotted onto the marked plate with the help of a capillary tube or pipette.

n-Hexane: Ethyl acted (5 ml: 5 ml) was used as mobile phase. The TLC plate was kept in a chromatographic chamber containing the respective solvent system and the chamber was covered with glass plate to prevent the evaporation of the solvent. The plate was allowed to remain in the chamber till the solvent reached up to 9 cm distance. The plate was then observed in UV-fluorescence analysis cabinet at short and long wavelengths.

TLC finger printing plate was derivatized with anisaldehyde sulphuric acid reagent followed by heating at 100 °C till coloured bands of various secondary metabolites appeared. The observations were taken before and after derivatization in visible as well as ultraviolet light.

RI (Retention factor) values were calculated as follows:

\[
RI = \frac{\text{Distance traveled by substance}}{\text{Distance traveled by solvent}}
\]

2.15. Column chromatography of partially purified alcohol fraction of *Thevetia peruviana* leaf extract

10 gm of dried and partially purified alcohol fraction of *Thevetia peruviana* leaf was dissolved in the mobile phase i.e. 5 ml n-Hexane and 5 ml Ethyl Acetate and this solution was subjected to column chromatography. Glass column (Merck: 100–200 mm) filled with 650 gm of silica gel was used for column chromatography. According to the colour band developed in column different fractions of extract containing various secondary metabolites were collected. All fractions obtained from column were dried in rotary vacuum evaporator under reduced pressure. These fractions were screened for their antifungal activity against *A. solani*. The fraction showing best antifungal activity against *A. solani* was subjected to further purification and characterization for active molecule via gas chromatography and mass spectrometry analysis [13,14].

2.16. Identification and structure determination of active molecule by gas chromatography/mass spectrometry (GC-MS)

The GC MS analysis was performed on a GC (PerkinElmer) system coupled to Perkin-Elmer Turbo Mass MS. HP1-MS capillary column (30 m × 0.25 μm × 0.25 μm) was used under the following conditions: oven temperature programmed from 70 °C for 10 min, then gradually increased at 290 °C at 3 min; injector temperature, 250 °C, carrier gas Helium, flow rate 1 ml/min; the volume of injected sample was 0.4 μl; split ratio 1:60; ionization energy 70eV: Run time 40 min. The relative amount of each component was calculated by comparing its average peak area to the total area. The identification of the separated volatile compounds was done through retention indices and mass spectrometry by comparing mass spectra of the unknown peaks with those stored in the Nist 98/Nbs 75 K GC-MS library.

3. Results and observations

In order to check antifungal activity of crude and partially purified extract, extracts were dissolved in inert solvent i.e. acetone. It was observed to be neutral for growth of test fungus. Results of antifungal activity of crude as well as partially purified extracts were given in Tables 1 and 2 respectively (Fig. 1 A, B). Inhibitory activity of standards and control tabulated in Table 3 (Fig. 1 C). In case of crude extract, 100% alcoholic extract was observed to be best which showed 71.48% inhibition followed by aqueous and 50% hydroalcoholic extract which showed inhibition percent of 56.60% and 20.24% respectively.

Among the partially purified fractions obtained, maximum mycelial growth inhibition of 74.38% was observed with partially purified alcohol fraction of *T. peruviana* leaf extract. The inhibitory effect of leaf extract of *Thevetia peruviana* on morphology and reproduction of test fungi are depicted in Tables 4 and 5 and Fig. 2 A, B. There was normal growth pattern i.e. abundant mycelial growth and sporulation of *A. solani* in control tube. Sequential decrease in conidial size and raise in mycelial width was observed with the increase in concentration of leaf extract. *A. solani* also showed decreased sporulation with rising concentrations of the extract. Average mycelium width in control was 1.35 μm, which increased up to 5.40 μm at 1.25 mg/ml 74.57% reduction in conidial size was observed at 1.25 mg/ml concentration.

Table 6 showed the results of phytochemical screening of *T. peruviana* leaf extract prepared in different polarity of solvents. In petroleum ether fraction, steroids and tannins were found. Benzene fraction of leaf extract contained tannins only. All flavonoids, steroids, carbohydrates, tannins and saponins were observed in methanol fraction. Saponins were found both in aqueous as well as chloroform extract and acetone fraction contained flavonoids and saponins.

RI values and different colours of bands observed after TLC of active fraction are seen in Table 7 and Fig. 3A, total 7 bands were observed after TLC of active fraction of *T. peruviana* leaf extract. Every band showed its characteristics RI value calculated using the formulae. Active fraction (alcoholic fraction) of leaf extract of *T. peruviana* was also subjected to column chromatography. In column fractionation of alcoholic leaf extract of *T. peruviana*, total 9 fractions were collected. Each of 9 fractions was further fractionized via TLC. Different colour bands from one to three were observed for different column fractions. The TLC plate was observed under UV light at 254 nm before and after derivatization (Fig. 3B). Table 7 shows the results of TLC of different column fractions. Different bands showed characteristics RI value and colour after derivatization using anis-aldehyde reagent on the TLC plates. RI value of...
different bands observed were ranges from 0.62 to 0.94.

GC-MS Analysis Fractions obtained from column chromatography was subjected to determine antifungal activity against plant pathogenic fungus *A. Solani*. Table 8 and Fig. 4 depicted the results of antifungal activity of column fractions. As compared to control, maximum inhibition in growth of test fungus was observed by fraction F9 i.e followed by F7, F3, F1, F5, F6, F8, F2 and F4. All column fractions were checked for antifungal activity against test pathogens. Among the fractions most significant inhibitory activity against *A. solani* was shown by fraction F9. This fraction F9 was subjected to GC-MS analysis for further refining and identification of active compound it contained. Fig. 5 showed the chromatogram obtained for fraction F9. There was total number of one compound seen on GC-MS chromatogram.

### 4. Discussion

Herbal renaissance ‘is happening all over the globe and people

| S. No | Column fraction | Growth Diameter after 7 days (mm) ± SD | % Mycelial growth inhibition |
|-------|-----------------|----------------------------------------|-----------------------------|
| 1.    | F1              | 40.66 ± 0.577                           | 49.59                       |
| 2.    | F2              | 53.66 ± 0.577                           | 33.47                       |
| 3.    | F3              | 39.33 ± 0.577                           | 51.23                       |
| 4.    | F4              | 56.00 ± 0.00                            | 30.57                       |
| 5.    | F5              | 41.66 ± 0.577                           | 48.35                       |
| 6.    | F6              | 45.66 ± 0.577                           | 43.39                       |
| 7.    | F7              | 34.66 ± 0.577                           | 57.02                       |
| 8.    | F8              | 50.66 ± 0.577                           | 37.19                       |
| 9.    | F9              | 20.66 ± 0.577                           | 74.38                       |
| 10.   | Control (Water) | 80.66 ± 0.577                           | NI                          |

Fig. 3. Thin Layer Chromatography (TLC) of various fractions of *Thevetia peruviana* leaf. A. (i) TLC of alcohol fraction of leaf under visible light (ii) TLC of alcohol fraction of leaf under UV light (after derivatization). B. (i) TLC of column fractions [1–9] of alcohol fraction of leaf under visible light (ii) TLC of column fractions of alcohol fraction of leaf under visible light (After derivatization).
returning to the naturals with hope of safety and security. Generally, the public is progressively moving towards acceptance and practice of herbal preparations [14]. In recent years, medicinal plants has started to increase its appeal to pharmaceutical companies and the scientific research community. This was due to evidences that these plant-derived compounds have a potential for many biological activities which include antimicrobial activity [15]. Scientists aim is to extract and characterize active phytocompounds found in plants, since they have provided high activity profile drugs [16].

Shade dried plant material is used for crude extraction as well as partial purification of extract. Most of the time, dried sample is favoured in view of the time required for experimental design. Sulaiman et al. [17] suggested that there should be a 3 h of time duration gap between harvest and experimental work in order to sustain freshness of samples, as fresh samples are delicate and be likely to deteriorate quicker than dried samples. According to Sulaiman et al. [18] maceration or percolation of fresh green plants or dried powdered plant material in water and organic solvent systems is best for extraction of plant material.

Results suggested that in the presence of leaf extract of *Thevetia peruviana*, reduction in conidial dimension and raise in mycelia width of *A. solani* was recorded. The manner of antifungal activity of leaf extract of *Thevetia peruviana* might be mainly due to cell wall attack and withdrawal of cytoplasm in the hyphae and eventually death of the mycelium [19]. Such modifications induced by the exposure to plant extract components may interfere with enzymatic reactions of wall synthesis and affect fungal morphogenesis and development.

Preliminary phytochemical studies and literature reveal that the leaves of *T. peruviana* contain iridoid glycosides, flavonoids [20], triterpenes [21], monoterpenes [22] and cardiac glycosides [23]. These compounds are known to cause structural and functional damage of cellular membrane. Impairment in membrane results in dissipation of two components of proton motive force, the pH gradients (ΔpH) and electrical potential (Δψ) [24,25]. Detrimental effects on proton motive force are strongly correlated with leakage of specific ions [26,27].

The medicinal plants are rich in secondary metabolites, a diverse group of chemicals, which include alkaloids, glycosides, amines, insecticides, steroids, flavonoids, and related metabolites, which have been extensively used in drug and pharmaceutical industry [28]. Flavonoids, tannins and phenolic acids belong to phenolic compound group of secondary metabolites. Phenolic compounds have been shown to exhibit antioxidant and antimicrobial activity [29]. In terms of antimicrobial properties, not the quantity of phenolic compounds and flavonoids is often important but the type of substances present in each extract [30]. Each fraction of plant extract prepared by successive extraction carries a specific set of secondary metabolites because solubility of secondary metabolites differs with different solvent. Qualitative phytochemical tests are used to detect phyto-constituents present in individual fraction. Several workers have studied phytochemical properties of plant extracts by qualitative phytochemical tests [31–33].

Active fraction with best inhibitory activity was incorporated to TLC and subsequently into column fractionation and GC-MS analysis to determine active ingredients present in extract. TLC is an analytical
Antimicrobial activity of plants extract may be attributed instead of existence of active ingredients in prepared infusions. Extraction processes usually used to extract active ingredients are crude extraction and partially purified extraction process [17,39,40]. The principle of series of extraction process is to isolate the soluble plant metabolites, and left behind the insoluble cellular mass (residue). Initial crude extraction procedure i.e. cold extraction of plant parts, able to extract compound mix of many plant metabolites, such as alkaloids, glycosides, phenolics, terpenoids and flavonoids.

GC-MS is the best technique to identify the bioactive constituents of long chain hydrocarbons, alcohols, acids, esters, alkaloids, steroids, amino and nitro compounds etc. Hence, Gas chromatography (GC) and Mass spectroscopy (MS) associated with particular detection techniques have become a sophisticated means for analysis of various active compounds from plant extract [41]. Priya et al. [42] Identified eighteen phytochemical constituents from the ethanolic extract of the leaves of Desmodium gyrans by Gas chromatogram Mass spectrometry (GC-MS). Nanadagopalan et al. [43] reported the presence of Phytol in the leaves of Kirigandula reticulata aerial parts, which was found to be effective in different stages of arthritis. Aja et al. [44] GC/MS analysis of Moringa oleifera leaf and seed which revealed that 9-octadecenoic acid (20.89%) constitutes the major constituent of the leaf extract while oleic acid (84%) is the major component of the seed extract.

Active compound, active column fraction (having best antifungal activity against A. solani) was subjected to GC-MS analysis. Only one active compounds observed on GC-MS chromatogram. According the peaks matching of GC-MS chromatogram with available data base showed the presence of benzoic acid in active fraction of Thevetia peruviana leaf extract which is already known chemical among the phytochemicals described for antimicrobial activity.

Benzoic acids are C6–C18aromatic carboxylic acids that serve as precursors for a wide variety of essential compounds and natural products [45]. Benzoic acid also provide skeleton for numerous specialized metabolites. Non-volatile, benzyl benzoil, and anthraniloyl compounds can also act as defense compounds [46]. Phytochemicals like benzoic acid have also been reported in extract prepared from plant parts of Caryophyllus aromaticus and Syzygium joabolanum by Ref. [47].

5. Conclusion

Plant-based formulations can successfully be used in agriculture to treat plant diseases and can limit the use of chemical control agents to safer levels. Thus use of plant extract against plant pathogenic fungus is an important field of study and will be a best alternative of existing chemical antifungal in the near future. Antifungal effect of plant extract is mainly attributed to existence of secondary metabolites. These active constituents binds to membrane receptors and alter the cascade action generated thus leading to cyto-morphological alterations in test pathogenic fungus and directly or indirectly effect the growth and reproduction of the pathogen, thus bringing about its inhibition. Phytochemicals responsible for antimicrobial activity can be extracted from plant parts by using solvents of different polarity. Partial purification leads partial fractionization of phyto-constituents (i.e. much in alcoholic fraction of Thevetia leaf extract) which can be further refined by column fractionation. This further refinement of phyto-constituents via GC-MS analysis is much sensitive and sophisticated technique of fractionation. In this study benzoic acid and its derivatives were observed in active column fraction which is already reported for antimicrobial activity in different plant.

Author statement

Bhanu Raj: Conceptualization, Methodology, Execute experiments, Software, Data curation, Writing- Original draft preparation. Sanjeev Meena: Methodology, Execute some experiments, Reviewing and Editing. Deepali Chittora: Writing- Reviewing and Editing Kanika
Sharma: Supervision.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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