**STAPHYLOCOCCUS SP. STRAIN MY 83295F: A POTENTIAL P,P'-DDT-DEGRADING BACTERIUM ISOLATED FROM PESTICIDE CONTAMINATED SOIL**

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**Received:** 26 August 2020; **Accepted:** 05 November 2020; **Published:** 30 December 2020

**Abstract:** Although DDT has been on the ban list by the Stockholm Convention for its environmental degradation, still a wave of emerging shreds of evidence has proved its circulation in developing countries. The intensity of environmental degradation and human health problems posed by residual DDT and its metabolites become of serious ecological concern, warranting a search for novel strains with a capacity to biodegrade these environmental contaminants. A new strain of the genus *Staphylococcus* was isolated from pesticide-contaminated soil. The 16S rRNA and phylogenetic analyses were used to identify the isolate and the 16S rRNA partial gene sequence was deposited in the NCBI GenBank as *Staphylococcus* sp. strain MY 83295F. The isolate was capable of growing in up to 60 mg L\(^{-1}\) of p,p’-DDT as the sole carbon source at an optimum pH of 6.5 and optimum temperature of 30°C within 120 h. Zn\(^{2+}\) has demonstrated a stimulatory effect on the growth of the strain in p,p’-DDT containing medium. However, Fe, Cu, Pb, Hg, Ag, and Cr ions showed inhibitory effects on the strain’s growth in the medium. The strain could be a handy tool for the bio-cleansing of residual p,p’-DDT in the contaminated environment.

**Keywords:** DDT biodegradation, *Staphylococcus*, 16S rRNA, pesticide contaminant, heavy metals.

**1. Introduction**

Among the most notorious and persistent agrochemical environmental contaminants, the dichlorodiphenyltrichloroethane (DDT) still occupies a prominent position in the ranking of the hazardous environmental pollutants. This arises due to its persistence and potential accumulation in both biotic and abiotic components of the ecosystem (Bussolaro et al., 2012; Devi, 2020). Although DDT and other Persistent Organic Pollutants (POPs) have been on the ban list by the Stockholm Convention for their environmental degradation capacity, DDT is, still in circulation in developing countries (Abdul Kader, 2019). A comprehensive report released by the United Nations Environmental Protection (UNEP) on the current status of DDT from 2015 to 2017 in developing countries indicated that India, Mozambique, South Africa, and Zimbabwe reported the use of DDT (UNEP, 2019). While Botswana, Eswatini, Ethiopia, Eritrea, Madagascar, Marshall Islands, Namibia, Uganda, Venezuela, and Zambia have refused to give any response on the status of the use of...
DDT between 2015-2017, despite all the efforts by the UNEP. This perhaps signals the use of DDT for vector control in these countries (UNEP, 2019).

It was documented that in Nigeria several tons of chlorinated pesticides are used every year for both agriculture and disease-borne control (Asogwa and Dongo, 2009), and consequently, remnants of DDT were recently traced in domestic water samples (Ogbeide et al., 2015). Many studies across the world have reported the traces of DDT in ambient air, domestic water, and various food sources (Bussolaro et al., 2012; Mendes et al., 2019; Thompson et al., 2019). The levels of DDT reported in many food items exceeded the tolerable daily limits stated by the U.S. Environmental Protection Agency (Sheldon et al., 2019).

DDT and its metabolites such as dichlorodiphenyldichloroethylene (DDE) have been implicated in many health-related problems. They were reported to be linked to endocrine disruption (Mnif et al., 2011; Piazza and Urbanetz, 2019). Truong et al. (2019) reported an association between DDT metabolites and long-term impairments of muscle health. Many studies have established the probable involvement of DDT in cancer induction (Hadara et al., 2016; Cohn et al., 2019). DDT was also documented to induce cognitive decline and alteration in maternal metabolomes (Medehouenou et al., 2019; Hu et al., 2019). Yu et al. (2019) demonstrated that DDT is associated with some risks of triggering apoptosis of skin fibroblast in some aquatic animals.

From the beginning of the last decade to date, many bacterial strains have been identified to degrade DDT and its metabolites; DDD and DDE (Fang et al., 2010; Hug et al., 2013; Wang et al., 2017; Pan et al., 2017; Xie et al., 2018). The microbial ability to degrade DDT have strengthened the hope of employing them in the bioremediation of DDT contaminated sites.

Although DDT degradation was reported to be a multistep process (Fang et al., 2010; Cutright and Erdem, 2012), it mostly occurs either through aerobic or anaerobic metabolism. With favourable conditions, suitable strains were shown to degrade DDT to 4-chlorobenzoic acid (4-CBA) aerobically or to 4,4-dichlorobenzophenone (4-DBP) under anaerobic conditions (Nadeau et al., 1998; Baczynski et al., 2010; Gao et al., 2011; Bao et al., 2012). *Alcaligenes* sp. and *Serratia marcescens* DT-1P were reported to degrade DDT to 4-CBA aerobically via the DDE metabolic pathway in the presence of additional carbon sources (Bidlan and Manonmani, 2002). However, DDT was shown to be rapidly degraded anaerobically to 4-DBP through the DDD reductive dechlorination pathway (You et al., 1996; Baczynski et al., 2010; Fang et al., 2010).

The intensity of environmental degradation and human health problems posed by DDT and its metabolites, particularly their persistent nature in the environment, searching a microbial community for novel strains with a capacity to bio-clean the environment is pertinent. This work was focused on the isolation and characterization of *p,p'-DDT*-degrading bacterium from the tropical contaminated soil.

2. Materials and Methods

2.1 Sample collection

A soil sample was collected from irrigation sites located at Phase I, Kadawa Irrigation Site, Hadejia-Jama’are River Basin, Kano State with a history of continued agrochemical farming activities for more than three decades. The sample was collected at the surface of the soil to the depths of 15 cm. The sampling was focused on these horizons because a large
portion of microbial activity occurs in these horizons. The soil sample was mixed evenly and 20 g was carefully put into a sterile container and taken to the laboratory at 4°C for bacterial isolation.

2.2 Preparation of media for bacterial growth

The Luria-Bertani medium (LB) was used for the bacterial growth.

2.3 Preparation of p,p’-DDT-minimal salt enrichment medium (MSM)

The Minimal Salt Medium (MSM) had the following composition as described by Pant et al. (2013) with some modifications: per litre of distilled water, 0.1 g CaCl₂.2H₂O, 0.08 g Ca(NO₃)₂ 4H₂O, 0.5 g MgCl₂.6H₂O, 1.0 g Na₂SO₄ and 1.0 g KH₂PO₄ were dissolved. Then before inoculation, MSM was enriched with p,p’-DDT (0.05 mg mL⁻¹). The p,p’-DDT-MSM contains p,p’-DDT as the only carbon source for the bacterial growth. Thus, growth in this media depends only on the strain’s ability to metabolize the p,p’-DDT pesticide.

2.4 Isolation of p,p’-DDT-degrading bacterium from soil samples

Isolation of strain MY 83295F was carried out using a modified isolation procedure described by Pant et al. (2013). Air-dried soil (0.5 g) was suspended in 20 mL of the prepared LB medium. The suspension was kept for 48 h at 30°C on a shaker. After the incubation, the LB medium was allowed to settle down for 2 h. An aliquot (150 μL) from the cleared LB supernatant was used to inoculate 6 mL of p,p’-DDT enrichment MSM. The culture was then incubated for 1 week at 30°C on a rotary shaker at 100 rpm. After incubation, 100 μL of the bacterial suspension was transferred into 4 mL of fresh p,p’-DDT enriched MSM and the incubation step was repeated. After four sequential cultivations, the isolate was inoculated on to MSM agar plates enriched with 0.05 mg mL⁻¹ of p,p’-DDT and incubated for 72 h at 30°C and the isolate formed was preserved. This ensures adequate exposure of the isolate to the p,p’-DDT as a sole carbon source.

2.5 Extraction of genomic DNA

A single loop of the isolate was used to inoculate 8 mL of LB medium. Followed by incubation at 37°C and 200 rpm for 24 h. The bacterial suspension (OD₆₀₀nm = 0.6) formed was centrifuged for 5 min at 10,000 rpm. Then the bacterial DNA was extracted following the protocol stated by Schmidt et al. (1991).

2.6 16S ribosomal RNA gene (16S rRNA) amplification

To amplify ~1.5 Kb gene from the isolated genomic DNA, 16S rRNA gene primers (BAC27F and BAC1492R) [16SRNA BAC27F: 5’-AGA GTT TGA TCC TGG CTC AAG-3’ and 16SRNA BAC1492R: 5’- GGT TAC CTT GTT ACG ACT T-3’] purchased from Sigma-Aldrich, United Kingdom, were used (Sangwan et al., 2005). The PCR was carried out using TC-E-48FA Gene Touch Thermocycler, Hangzhou Bioer Technology, (China). The total reaction volume was 15 μL, in which the reaction mix comprises of 1 μL of the genomic DNA, 1.5 μL of 10X TaqA buffer, 0.5 μL of each of 10 μM forward and reverse primers, 0.75 μL of 1.25 mM of MgCl₂, 0.15 μL of 0.25 mM of dNTP and 0.12 μL of Taq DNA polymerase in ddH₂O. The PCR protocol was set as follows: the initial melting temperature was 95°C for 5 min, 35 cycles each at melting temperature of 94°C for 0.5 min. The annealing temperature was 52°C for 0.5 min and extension at 72°C for 1 min. The final elongation was set for 10 min at 72°C.

After the final elongation cycle, the size of the DNA fragment was compared with the Hyper Ladder-1K marker Bioline (Lot No: H4-
Then 3 of the µL PCR product was mixed with 5XDNA loading buffer blue (1.5 µL) Bioline (Lot No: hLBB-415704) and loaded onto 1.5% agarose gel electrophoresis that has been stained with ethidium bromide. The electrophoresis was run for 35 min under 120V and 300mA current. The product was then visualized with the Syngene Gel Documentation System of Ingenius, England (IG31459). The presence of a product of the expected size was considered to be a positive result.

### 2.7 Agarose gel purification and sequencing of amplified 16S rRNA gene

The gel was purified using the PrepEase gel purification kit (Affymetrix inc., USA) by following the manufacturer’s protocol. The gel-purified product was sequenced using the protocols described by Sanger et al. (1977). Then, DNA sequence alignment was carried out using the ClustalW 2.0.12 version (http://www.clustal.org/). The sequence was then blasted in the National Center for Biotechnology Information (NCBI) nucleotide databases to identify the organism. The sequence was deposited in the NCBI GenBank under the accession number MN812290.

### 2.8 Phylogenetic analysis of the isolates

Phylogeny and evolutionary history of strain MY 83295F were constructed using the Neighbor-Joining method. The evolutionary distances were computed using the Maximum Composite Likelihood method (Tamura et al., 2013). The phylogenetic tree was constructed using MEGA version 6 software program.

### 2.9 Characterization of the isolate in minimal salt-\(p,p'\)-DDT enrichment medium

During the characterization of the strain MY 83295F in the \(p,p'\)-DDT enrichment medium, we have determined the optimum \(p,p'\)-DDT concentration (as a sole carbon source), pH, temperature and incubation time. The characterization was performed by modifications of the methods described by Mwangi et al. (2010) and Pant et al. (2013). The isolate’s capacity to grow in the \(p,p'\)-DDT enrichment medium was determined \textit{in vitro} after adjusting the \(OD_{600\text{nm}}\) of cell density of the inoculum source to 0.6. The inoculum (150 µL) was then inoculated into 4 mL of MSM media containing varying concentrations \(p,p'\)-DDT (10, 20, 30, 40, 50, 60 and 70 mgL\(^{-1}\)) at various pH values (5.5, 6.0, 6.5, 7.0, 7.5, 8.0 and 8.5) and incubated at different temperatures (20, 25, 30, 35, 40 and 45°C) under shaking (150 rpm) at different incubation periods of 24, 48, 72, 96, 120, 144 and 168 h. The experiments were conducted one factor at a time in triplicate.

### 2.10 Effects of heavy metals on \(p,p'\)-DDT utilization capacity of the isolates

The effect of each heavy metals (Fe, Zn, Cu, Pb, Hg, Ag and Cr) on \(p,p'\)-DDT degrading capacity of the strain MY 83295F was determined \textit{in vitro} after adjusting the \(OD_{600\text{nm}}\) of cell density of the inoculum source to 0.6, the cells (100 µL) were then inoculated into MSM-DDT enrichment media containing varying metal concentrations (0.2, 0.4, 0.6, 0.8 and 1.0 mgL\(^{-1}\)) and incubated on a rotary shaker (150 rpm) at 30 °C and pH 6.5 for 168 h. This protocol is a modified version of the procedure described by Sandrin and Maier, (2003).

### 3. Results and discussions

In this study, minimal salt-\(p,p'\)-DDT enrichment medium was used for the isolation and screening of \(p,p'\)-DDT bio-degrader bacterial strain that used \(p,p'\)-DDT as the sole carbon and energy source obtained from pesticide-contaminated agricultural soil. The biodegradation capacity of \(p,p'\)-DDT by the isolate was indicated by the formation of
turbidity as the index of the biomass formed in the minimal salt-\textit{p,p'-}DDT enrichment medium (Mwangi et al., 2010; Pant et al., 2013).

\textit{Staphylococcus} sp. strain MY 83295F was found to be a Gram-positive, non-spore-forming, non-motile, catalase-positive, urease, and cytochrome oxidase negative cocci. However, the strain was found to ferment D-glucose, starch and D-mannitol while negative with xylose and indole (data not presented here). A substantial literature on the phenotypic and biochemical characteristics of the genus \textit{Staphylococcus} showed clear similarities with the above-mentioned characteristics of strain MY 83295F (Bascomb and Manafi, 1998; Khattak et al., 2015; Karmaker et al., 2016).

The 16S rRNA gene sequencing was found to maintain reasonable accuracy and reliability for bacterial identification (Roy et al., 2013; Hong and Farrence, 2015). Thus, for the molecular identification of this isolate to confirm the preceding phenotypic and biochemical identifications, the 16S rRNA gene was amplified. The amplification product for the isolate revealed about 1500 bp upon running on 1.5\% agarose gel electrophoresis (Fig. 1.). Many researchers have earlier reported similar ranges of 16S rRNA gene amplification products between the ranges of 1200-1500 bp for the genera \textit{Staphylococcus} depending on the species, segment amplified or the type of primers used (Saruta et al., 1997; Jill and Clarridge, 2004; Mitra and Roy, 2010).

The 16S rRNA gene amplicon was successfully sequenced, revealing 1057 bp as the partial gene sequence for the strain MY 83295F. The sequence was blasted in the NCBI GenBank that revealed the genus of the strain as \textit{Staphylococcus}. The 16S RNA gene partial sequence was deposited in public databases of the National Center for Biotechnology Information (NCBI) GenBank as \textit{Staphylococcus} sp. strain MY 83295F under the universal accession number MN812290.

The BLAST search on strain MY 83295F showed about 80\% of the first hundred representatives were of the genus \textit{Staphylococcus}, revealing the highest sequence similarity of 99.53\% with \textit{Staphylococcus hominis} subsp. novobiosepticus strain GTC 1228, followed by \textit{Staphylococcus hominis} strain DM 122 with 99.15\%.

\begin{center}
\textbf{Fig. 1.} Agarose gel electrophoretic image of the 16S rRNA amplicon of MY 83295F strain. The gene was amplified using BAC27F and BAC1492R as forward and reverse primers respectively. L represents Hyper Ladder-1K marker Bioline (Lot No: H4-q111B)
\end{center}
The least % sequence similarity of 95.27% was found in *Staphylococcus agenesis* strain M4S-6 among the representatives of the genus *Staphylococcus*. The phylogeny and evolutionary analysis revealed a single clade of *Staphylococcus hominis* strain DM 122 and a cluster of strain MY 83295F and *Staphylococcus hominis* subsp. *novobiosepticus* strain GTC 1228 with a bootstrap value of 85%, indicating a closer relatedness of these organisms. However, the cluster of strain MY 83295F and *Staphylococcus hominis* subsp. *novobiosepticus* strain GTC 1228 indicated a bootstrap value of 59% (**Fig. 2**). This value is very low to state with certainty that strain MY 83295F belongs to the same subspecies with *Staphylococcus hominis* subsp. *novobiosepticus* strain GTC 1228. Perhaps, suggesting a new subspecies within the group of *Staphylococcus hominis*. Therefore, strain MY 83295F was tentatively designated as *Staphylococcus* sp. strain MY 83295F, where the sp. indicates an unclassified species of this strain, subject to in-depth taxonomical approach. Several studies on the identification and taxonomy of the genus *Staphylococcus* have reported a similar approach in the identification and grouping of *Staphylococcal* species (Stackebrandt and Goebel, 1994; Takahashi et al., 1999; Ghebremedhin et al., 2008; Naushad et al., 2016).

Microorganisms are vital tools for the removal of various toxic contaminants including the persistent chlorinated pollutants such as DDT from the environment (Reineke et al., 2011). Biodegradation of DDT by bacteria has been well documented, and DDT-degraders have been isolated (Mwangi et al., 2010; Pan et al., 2016). Strain MY 83295F was found to biodegrade and utilize *p,p*-DDT as sole carbon and energy source. Though some bacterial species were reported to tolerate < 20 mgL\(^{-1}\) of *p,p*-DDT as sole carbon and energy (Pant et al., 2013; Pan et al., 2016), strain MY 83295F, however, demonstrated higher tolerance and utilization capacity of up to 60 mg L\(^{-1}\) of *p,p*-DDT as the sole carbon source under aerobic condition (**Fig. 3a**).

**Fig. 2.** Phylogenetic and Evolutionary relationships of taxa of strain MY 83295F. The strain’s evolutionary position was indicated in a rectangular box. The evolutionary history was inferred using the Neighbor-Joining method.
Fig. 3. Effects of \( p,p' \)-DDT concentration (a) and incubation time (b) on the growth of strain MY 83295F in \( p,p' \)-DDT enrichment medium. For determination of the effect of the incubation time on the growth of the strain, 60 mgL\(^{-1}\) of \( p,p' \)-DDT was used. The turbidity of the medium is an index of growth of the isolate in the \( p,p' \)-DDT enrichment medium, which was determined spectrophotometrically as optical density (OD) at 600 nm. The experiments were conducted in triplicate.

Some isolates were also documented to tolerate up to 50 mgL\(^{-1}\) of DDT when other carbon sources were supplemented (Kantachote et al., 2003; Barragan-Huerta et al., 2007; Fang et al., 2010).

The ability of a microorganism to depend on DDT as a carbon source depends on the organism’s capacity to mineralize the DDT and obtain energy from the process (Fang et al., 2010; Pan et al., 2016). Strains MY 83295F demonstrated longer lag phases of nearly 48 h for the initial DDT degradation (Figure 3b). The delay observed in the initial rate of DDT degradation in the strain could be attributed to the delay in the production of enzyme machinery for the degradation. However, the strain was able to moderately biodegraded and utilized \( p,p' \)-DDT, precipitating total biomass of 0.130 (OD\(_{600nm}\)) in 120 h (Fig. 3b).

As a complex process, bacterial DDT mineralization is largely influenced by some environmental determinants such as pH, temperature and DDT concentration (Aislabie et al., 1997; Bidlam and Manonmani, 2002). Optimization of these parameters is therefore critical for the application of a microbial entity for the DDT biodegradation. Strain MY 83295F demonstrated a mesophilic behaviour by exhibiting \( p,p' \)-DDT degradation within a wide range of temperatures between 20-45°C. However, strain MY 83295F showed an optimum temperature of 30°C (Fig. 4a). This is not surprising by considering the environmental conditions of the tropical region where this strain was isolated. A fluctuation in temperature, either below or above the optimum value, the strains’ DDT degradation capacity is lowered.

Strain MY 83295F showed growth capacity in \( p,p' \)-DDT enrichment media in both slightly acidic and alkaline conditions, with initial pH ranging from 5.5 to 7.5. However, the strain demonstrated an optimum pH of 6.5 (Fig. 4b). A substantial literature has reported an optimal bacterial DDT degradation within a range of pH close to neutral and temperature range around 30°C that correspond to those shown by this strain (Mwangi et al., 2010; Fang et al., 2010; Pan et al., 2016; Raju and Bidlan, 2018).
In biological systems, including microorganisms, pH plays a significant role in changing the ionic character of the constituent amino acids in enzymes and other intracellular and membrane proteins. This perhaps, influences DDT degradation capacity of this bacterial strain, either by affecting the DDT membrane transport system or degradation enzymes.

In most cases, organic pollutants such as DDT and heavy metals co-contaminate the environment, and more or less, the later might influence the degradation rate of the former as reported by Lovecke et al. (2015). The effects of heavy metals on DDT degradation capacity of strain MY 83295F showed both stimulatory and inhibitory effects. Only Zn$^{2+}$ at 0.2 mgL$^{-1}$ has demonstrated enhancement effect on p,p$'$-DDT degradation in Staphylococcus sp. strain MY 83295F, precipitating an increase in the biomass in p,p$'$-DDT enrichment medium after 168 h. An increase in the Zn$^{2+}$ concentration above 0.2 mgL$^{-1}$ showed a dramatic decline in p,p$'$-DDT degradation, and reduction in the total biomass of the strain MY 83295F (Fig. 5).

Zn$^{2+}$ is physiologically essential for bacterial growth. It serves as a cofactor for many microbial metalloenzymes and other structural and regulatory functions. Bacteria have systems called cation diffusion facilitator (CDF) proteins that modulate their survival in the Zn$^{2+}$ contaminated environment (Guffanti et al., 2012). These bacterial proteins are responsible for Zn$^{2+}$ homeostasis via Zn$^{2+}$-uptake/import and Zn$^{2+}$-efflux/export mechanisms (Suryawati, 2018). The genus Staphylococcus was reported to have these CDF proteins (Nies, 2003).

Thus, Staphylococcus sp. strain MY 83295F might have very active Zn$^{2+}$ homeostasis systems that warrant growth enhancement in the Zn$^{2+}$-p,p$'$-DDT enrichment medium. Indeed, the DDT degradation enzyme machinery might interact with the divalent ion such as zinc as presented by Mansouri et al (2017), leading to the enhancement of catalytic ability by the enzyme machinery.

A pattern of inhibition demonstrated by Fe$^{2+}$, Cu$^{2+}$ (Fig. 6a and b) was more favourable to the growth of the strain 83295F relative to that of Pb$^{2+}$(Fig. 7a), Hg$^{2+}$ (Fig. 7b), Ag$^{2+}$ (Fig. 8a) and Cr$^{2+}$ (Fig. 8b). This could be linked to the essentiality of Cu$^{2+}$ for some metabolic processes in bacteria. However, it has been observed that presence of both organic and metal pollutants resulted in metal toxicity in
bacteria, mostly by interacting and inhibiting the bacterial enzymes and thus, inhibiting organic pollutant biodegradation (Angle and Chaney, 1989; Sandrin and Maier, 2003; Murata et al., 2005). Also, metal oxyanions, such as chromate, mimic the structure of essential non-metal oxyanions, such as sulfate, and interfere with their biological functions (Sandrin and Maier, 2003). Furthermore, mercuric and silver cations form strong toxic complexes which make them dangerous for any physiological functions, in addition to their inhibitory binding to the SH group of the variety of bacterial proteins (Nies, 1999). Metal ions generally affect organic biodegradation by altering both the physiology and ecology of the organic bio-degraders (Sandrin and Maier, 2003). The net effect of exposure to Cu\(^{2+}\), Pb\(^{2+}\), Hg\(^{2+}\), Ag\(^{2+}\) and Cr\(^{2+}\) on the \(p,p'\)-DDT degradation by strain MY 83295F was thus, reduced biodegradation rates and failure to effectively degrade the \(p,p'\)-DDT.

**Fig. 5.** Effect of Zn\(^{2+}\) concentration on the growth of strain MY 83295F in \(p,p'\)-DDT (60 mgL\(^{-1}\)) enrichment medium. The turbidity of the medium is an index of growth of the isolate in the \(p,p'\)-DDT enrichment medium, which was determined spectrophotometrically as optical density (OD) at 600\(_{\text{nm}}\). The experiments were conducted in triplicate.

**Fig. 6.** Effect of Fe\(^{2+}\) (a) and Cu\(^{2+}\) (b) concentrations on the growth of strain MY 83295F in \(p,p'\)-DDT(60 mgL\(^{-1}\)) enrichment medium. The turbidity of the medium is an index of growth of the isolate in the \(p,p'\)-DDT enrichment medium, which was determined spectrophotometrically as optical density (OD) at 600\(_{\text{nm}}\). The experiments were conducted in triplicate.
Fig. 7. Effect of Pb$^{2+}$ (a) and Hg$^{2+}$ (b) concentrations on the growth of strain MY 83295F in $p,p'$-DDT(60 mg L$^{-1}$) enrichment medium. The turbidity of the medium is an index of growth of the isolate in the $p,p'$-DDT enrichment medium, which was determined spectrophotometrically as optical density (OD) at 600 nm. The experiments were conducted in triplicate.

Fig. 8. Effect of Ag$^{2+}$ (a) and Cr$^{2+}$ (b) concentrations on the growth of strain MY 83295F in $p,p'$-DDT(60 mg L$^{-1}$) enrichment medium. The turbidity of the medium is an index of growth of the isolate in the $p,p'$-DDT enrichment medium, which was determined spectrophotometrically as optical density (OD) at 600 nm. The experiments were conducted in triplicate.

**Conclusions**

A strain MY 83295F of the genus *Staphylococcus* was isolated from pesticide-contaminated soil. The strain was capable of growing in up to 60 mg L$^{-1}$ of $p,p'$-DDT as the sole carbon and energy source at an optimum pH of 6.5 and optimum temperature of 30 °C within 120 h. Zn$^{2+}$ demonstrated a stimulatory effect on the growth of the strain in the $p,p'$-DDT enrichment medium. However, Fe, Cu, Pb, Hg, Ag and Cr ions showed various patterns of an inhibitory effect on the growth of the strain. Thus, a simultaneous incidence of $p,p'$-DDT and the inhibitory heavy metals in same environment may alter the $p,p'$-DDT biodegradation potentiality of the strain. This strain could be a handy tool for the bioremediation of residual $p,p'$-DDT contaminant.
Conflict of Interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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