Diversity and Chemical Composition of Weeds in Sand-Filled Mangrove Forest at Eagle Island, Niger Delta, Nigeria

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Abstract

Mangroves are habitat specific and grow mainly in swampy soil, but due to anthropogenic activities (e.g. sand mining) other species had encroached into their habitat. It is thus hypothesized that change in species diversity will lead to change in soil chemistry. In a 40 m × 90 m plot, diversity index ($H$) and importance value ($I_v$) of weed were estimated. Soil and weed samples were collected and analyzed for total hydrocarbon content (THC), Zinc (Zn), Lead (Pb) and Cadmium (Cd). All samples were analyzed with atomic absorption spectrophotometric method using the HACH DR 890 calimeter (wavelength 420 nm). The result shows that swampy soils were more acidic (3.1 - 3.5) than sandy soils (4.2 - 4.7). Swampy soil was also more saline and thus has higher conductivity (8320 - 9880 µS/cm) than sandy soil (4320 - 5650 µS/cm). Mangrove swamp had higher total organic carbon (TOC) (2.25% - 3.41%) than sandy soil (0.12% - 0.21%). There was a significant difference in THC and heavy metals in soil ($F_{8,63} = 2.04, P < 0.05$), but there was no significant difference in THC and heavy metals in plant species ($F_{8,63} = 247.0, P > 0.05$). Concentration of THC and heavy metal was higher in plant than in soil. Reissantia indica, an aquatic weed, had the highest concentration of THC in root soil. A total of fifteen (15) weed species were identified, out of which Mariscus longibracteatus had the highest diversity ($-0.366$) followed by Mariscus ligularis ($-0.339$) and Paspalum vaginatum ($-0.270$). Similarly, M. longibracteatus had the highest importance value in the study site ($I_v = 58.24$). This result implies that the presence of weed species in mangrove forest is an indicator of human disturbance of the ecosystem. It also means that the weeds were bioaccumulating THC and heavy metals present in the soil.
1. Introduction

Mangroves are habitat specific and inhabit swampy and saline environment [1] [2] [3] [4]. Mangrove weed are plants that are found in disturbed mangrove forest such as reclaimed land, sand filled and dredged sites [5]. Weeds are unwanted plants that grow in any place that is favorable for their growth and survival such as water, soil, tree trunk, wall of building and coastal soil [6]. Aquatic weeds grow in water and have effects on coastal environment [7]. For instance these weeds can change the ecology of the area by supplying or utilizing soil nutrients, which may be detrimental to the native species [8] [9]. Weed can affect fish spawning capability by obstructing breeding grounds [10]; they can also increase the heavy metal concentration through the actions of their root, which break up the parent soil to expose more metals. Humans also introduce seeds of weed into the mangrove forests, which embed in the soil and grow. Growth of weed in disturbed mangrove forests converts indigenous mangrove soil to sandy soil. The weed species are able to grow at the fringes and perimeters of the mangrove forest that had been cut down for the purpose of creating access way for pipelines [11]. This is because the pipeline route is usually covered with sandy soils brought elsewhere, which further introduce and accelerate weed growth [12]. Furthermore, the sand filling of mangrove forest also changes the soil chemistry when muddy soils are converted to sandy soil. Change in soil chemistry can lead to increased heavy metal load, which has ripple effect on the aquatic environment [13]. For instance, fishes and other aquatic organisms bioaccumulate the excess heavy metals, which they pass on to humans who feed on them [14] [15]. Over the years several questions had been asked as to the role played by the mangrove weeds, which are often found in degraded mangrove forests. Do these foreign plants growing in mangrove forest play positive and/or negative roles? These questions are yet to be addressed. However, it is revealed by previous studies that in agricultural farms weeds can act as habitat for insect pollinators e.g. butterfly, grasshoppers, praying mantis beetles etc., which is a positive role for nearby plants. Nevertheless, weeds can play negative role by acting as nesting sites for pathogenic organisms such as mosquitoes, black fly etc. [16]. The growth of the weeds near the mangrove forest can also bring in destructive herbivores such as locust, caterpillars and insect pests that feed on leaves. In the Niger Delta there are limited studies to address these questions. Therefore, this study is aimed at investigating the diversity of weed and other plant species found in sand filled mangrove forests, and to determine the concentration of some heavy metals in soil and plant. The objectives of this study are: 1) to determine the diversity index ($H$) and importance value ($I_v$) of the

Keywords

Heavy Metals, Mangrove Weed, Soil, Species Diversity, Sand Fill, Swamp
weed species 2) to determine the THC and heavy metal concentration in both soil and plant, and 3) to compare the THC and heavy metal concentration in plant and soil.

2. Materials and Methods

2.1. Description of Study Area

The study area is an abandoned sand dump located at Eagle Island (N 4°47.317' and E 6°58.593'), directly behind the Rivers State University (Figure 1). The area has a warm and humid climate with two seasons, dry and wet seasons [17]. The soil of the area is between sandy to muddy, and also whitish to dark brown in color [18]. The area was once a mangrove forest, but was cut down to make way for sand mining five years ago. At the end of the sand mining activity the area was later abandoned to its fate. The company that operated the sand mine left the site with heaps of white sand still on the ground surface [19]. This abandoned sand became a platform on which a variety of weed and other plant species grow over the years. These species are plants that naturally cannot grow in swampy soil but because of the conversion of the sand from swampy to sandy soil weeds do proliferate. Moreover, the site is surrounded by a river channel which brings in sediments and seeds of plants during high tide. At the edges of the sand dump are heaps of mangrove soils placed to prevent the entry of water during high tide.

2.2. Sample Collection

In an area measuring 3600 m² (i.e. 40 m × 90 m) eight plots were delineated from where weed and soil samples were randomly collected (Figure 2). The plants
Figure 2. The four most dominant plant species growing in sand filled mangrove forest at Eagle Island, Niger Delta (a) *Mariscus ligularis* (b) *Paspalum vaginatum* (c) *Reissantia indica* (d) *Acrostichum aureum*.

were counted and each weed samples collected and placed in a cellophane bag. Similarly soil samples were collected beneath the plant root 5 cm below the soil with a soil augur and also placed in a cellophane bag. All the samples were put in a cooler and transported to the laboratory for physico-chemical analysis.

2.3. Physico-Chemical Analysis

In order to determine the soil chemistry of the study area, physico-chemical analysis of soil were done for the following parameters: pH, conductivity, total hydrocarbon (THC), total organic content (TOC), Nickel (Ni), Lead (Pb), Chromium (Cr), Magnesium (Mg), Potassium (K), Sodium (Na), Calcium and (Ca), while for the plant sample the following were parameters were analyzed: Cd, Pb, Zn and THC using standard laboratory procedures described below.

2.4. Procedures of THC Analysis

It involved the use of spectrophotometric method using the HACH DR 890 calorimeter (wavelength 420 nm). The samples were crushed and 2 g of the crushed sample was weighed into a glass beaker and 20 ml of hexane was added, and with the aid of a glass rod, the mixture was homogenized by stirring. Afterwards, the sample was filtered in a glass funnel packed with cotton wool, silica gel and anhydrous sodium sulphate. After this, 10 ml of the filtered organic extract was transferred into a 10 ml sample curvet and inserted into the calorimeter. The detection limit for THC is 0.01 mg/l. The above method followed the procedures of [20].
2.5. Procedures of Heavy Metal Analysis

Heavy metal extraction followed the example of [21]. Aliquots of 0.25 g of air dried sediment samples were weighed into a Teflon inset of a microwave digestion vessel and 2 ml concentrated (90%) nitric acid (Sigma-Aldrich, Dorset, UK) were added. The metals were extracted using a microwave accelerated reaction system (MARS Xpress, CEM Corporation, Matthews, North Carolina) at 1500 W power (100%), ramped to 175°C in 5.5 min, held for 4.5 min, and allowed to cool down for 1 h. The cool digest solution was filtered through the Whatman 42 filter paper and made up to 100 ml in a volumetric flask by adding de-ionized water. All chemicals and reagents used were of analytical grade and of highest purity possible. Analytical blanks were prepared with each batch of the digestion set and analyzed (one blank for every set of 6 samples) in the same way as the samples. The detection limit for the three metals analyzed in mg/l i.e. Zinc, Cadmium and Lead is 0.001, 0.001 and 0.002 respectively.

2.6. Procedures of Physico-Chemical Analysis

The pH and conductivity were measured using Sper Scientific 86003A multiparameter meters with probes calibrated with standard solutions. Total Organic Carbon (TOC) was determined using Walkey-Black titrimetric method.

2.7. Identification of Weed Species

The weeds collected from the site were sent to the laboratory and identified using a handbook of West African weed by [7]. The weeds were identified based on the shape of the leaves, color, seeds, inflorescence and fruit produced.

2.8. Statistical Analysis

An analysis of variance (ANOVA) was conducted to determine whether there was a significant difference in THC and heavy metal concentration in soil and plant [22]. Bar graphs were then used to illustrate the significance and difference in concentration in plant and soil. All analyses were done in [23].

2.9. Data Analysis

The stand basal area, G, measured in m²/ha, (Equation (1)) refers to the summation of all individual basal areas per unit ground area [24]. Where, \( g_i \) (m²) is the basal area of a single stand (Equation (2)), while 25 (400) is the plot size of the 40 m × 90 m (3600 m²) main plot, and a 5 m × 5 m (25 m²) sub-plot, as the conversion factor to 1 hectare [25].

\[
G = \sum_{i=1}^{N} g_i \times 25(3600) \tag{1}
\]

\[
g = \pi \times \left( \frac{dbh}{2} \right)^2 \tag{2}
\]

The importance value, \( I_i \), is a quantitative parameter used to show the significance of each species within a stand. It is a summation of density, frequency
and dominance (Equations (3), (4), (5) and (6)). The importance value \( I_v \) was calculated using the formula of [24].

\[
I_v = \text{Relative Density} \pm \text{Relative Frequency} \pm \text{Relative Dominance} \tag{3}
\]

\[
\text{Relative Density} \left( \% \right) = \frac{\text{no of individuals of species} \left( \frac{N}{\text{ha}} \right)}{\text{total number of individuals} \left( \frac{N}{\text{ha}} \right)} \times 100 \tag{4}
\]

\[
\text{Relative Frequency} \left( \% \right) = \frac{\text{frequency of species} \left( n \right)}{\text{total number of species} \left( \frac{N}{\text{ha}} \right)} \times 100 \tag{5}
\]

\[
\text{Relative Dominance} \left( \% \right) = \frac{\text{total basal area of species}}{\text{Basal area of all species} \left( G \right)} \times 100 \tag{6}
\]

To determine the species diversity, the Shannon Index \( H \) was used (Equation (7)), and is based on natural logarithm which considers low and high diversity species based on abundance of species. Higher values of diversity index imply higher biodiversity of that plant species and vice versa [26]. Species diversity is used for this study because it serves as an indicator of human disturbances [27].

\[
H = \sum_{i=1}^{1} p_i \ln \left( p_i \right) \tag{7}
\]

where,

\( H = \) Shannon diversity index;

\( \sum_{i=1}^{1} = \) summation;

\( \ln = \) natural logarithm.

### 3. Results

#### 3.1. Physico-Chemical Analysis

Result (Table 1) of the physico-chemistry of the study area shows that swampy soils are more acidic (3.10 - 3.50) than sandy soil (4.20 - 4.70) that harbor the weed species. Swampy soils are also more saline and thus have higher ability to conduct electrons (8320.00 - 9880.00 µS/cm) as compared to sandy soil (4320.00 - 5650.00 µS/cm). Mangrove swamp has higher total organic carbon (TOC) (2.25% - 3.41%) than sandy soil (0.12% - 0.21%). Furthermore, swampy soil has more concentration of heavy metals i.e. Ni, Pb and Cr. In addition, swampy soils are also high in K and Na [28]. In contrast, sandy soil had more Mg content.

| Sample Identity | pH   | Conductivity (µS/cm) | THC (mg/kg) | TOC (%) | Ni (mg/kg) | Pb (mg/kg) | Cr (mg/kg) | Mg (mg/kg) | K (mg/kg) | Na (mg/kg) | Ca (mg/kg) |
|-----------------|------|----------------------|-------------|---------|------------|------------|------------|------------|----------|-----------|-----------|
| Swamp 1         | 3.10 | 8320.00              | 5.63        | 3.41    | 5.66       | 5.60       | 0.59       | 134.50     | 14.20    | 2389.00   | 4.20      |
| Swamp 2         | 3.50 | 9880.00              | 2.33        | 2.25    | 6.35       | 7.27       | 7.37       | 62.20      | 192.00   | 3528.00   | 18.90     |
| Sand 1          | 4.70 | 4320.00              | 5.50        | 0.12    | 1.04       | 0.41       | <0.001     | 315.00     | 10.00    | 625.70    | 82.10     |
| Sand 2          | 4.20 | 5650.00              | 7.84        | 0.21    | 1.19       | 1.78       | 44.87      | 306.10     | 11.90    | 351.60    | 1.60      |
3.2. Concentration of THC and Heavy Metals in Soils Collected from Root of Weed Species

The results for the nine most dominant weed species (Table 2, Figure 3) indicate that there is a significant difference in THC and heavy metal concentration in soil ($F_{8,63} = 2.04, P < 0.05$). *R. indica* has the highest concentration of THC in root soil. There was also a significant difference in the concentration among the parameters *i.e.* THC and heavy metals ($P < 0.05$). Zn and THC had the highest soil concentrations (Table 2). *M. longibracteatus* had the highest Zn ($7.67 \pm 0.005$ mg/kg) followed by *A. areum* ($5.79 \pm 0.01$) and *M. ligularis* ($5.60 \pm 0.01$).

![Figure 3. Mean THC and heavy metal bioaccumulation in (a) soil and (b) plant samples collected in sand filled mangrove forest at Eagle Island, Niger Delta, Nigeria.](image)

| Species          | Metals mg/kg          |
|------------------|-----------------------|
|                  | Cadmium | Lead | Zinc   | THC     |
| *A. areum*       | 0.002 ± 0.0004 | 0.002 ± 0.0005 | 5.79 ± 0.01 | 5.59 ± 0.04 |
| *C. longipinna*  | 0.002 ± 0.0005 | 0.002 ± 0.0005 | 2.97 ± 0.01 | 2.31 ± 0.03 |
| *C. mindorensis* | 0.002 ± 0.0004 | 0.001 ± 0.0000 | 1.12 ± 0.01 | 7.35 ± 0.05 |
| *D. oliveri*     | 0.002 ± 0.0000 | 0.002 ± 0.0005 | 1.54 ± 0.06 | 5.46 ± 0.04 |
| *M. ligularis*   | 0.002 ± 0.0005 | 0.001 ± 0.0000 | 5.60 ± 0.01 | 8.79 ± 0.01 |
| *M. longibracteatus* | 0.001 ± 0.0000 | 0.001 ± 0.0000 | 7.67 ± 0.01 | 7.55 ± 0.05 |
| *P. vaginatum*   | 0.002 ± 0.0005 | 0.002 ± 0.0005 | 1.94 ± 0.01 | 8.84 ± 0.06 |
| *R. indica*      | 0.001 ± 0.0000 | 0.001 ± 0.0000 | 1.99 ± 0.03 | 137.1 ± 1.30 |
| *S. indica*      | 0.002 ± 0.0005 | 0.002 ± 0.0005 | 1.99 ± 0.01 | 7.75 ± 0.09 |
mg/kg). \textit{R. indica} had the overall highest THC concentration (137.1 ± 1.30 mg/kg) followed by \textit{P. vaginatum} (8.84 ± 0.06 mg/kg) and \textit{M. ligularis} (8.79 ± 0.01 mg/kg). Cd and Pb range between 0.001 - 0.002 mg/kg. The order of importance for the chemicals in soils is THC > Zn > Cd > Pb.

### 3.3. Concentration of THC and Heavy Metal in Weed Species

In contrast, there was no significant difference in THC and heavy metal concentration in the body of the plants ($F_{8,63} = 247.0$, $P > 0.05$). However, there was a significant difference in the concentration of THC and heavy metals ($P < 0.05$). Zn and THC had the highest concentration in weed (Table 3). \textit{A. areum} had the highest Zn concentration (52.67 ± 0.45 mg/kg) while \textit{M. ligularis} has the highest THC. Cadmium had the least concentration in weed. The order of importance in weed is Zn > THC > Pb > Cd.

### 3.4. Diversity of Weeds in Sand-Filled Mangrove Forests

A total of fifteen weed species were recorded within the sand filled area of the mangrove forest (Table 4). Most of the species fall under aquatic low land and dry land weeds, which are dicotyledons and monocotyledons [2]. The result indicate that \textit{Mariscus longibracteatus} had the highest diversity (−0.366) followed by \textit{Mariscus ligularis} (−0.339) and \textit{Paspalum vaginatum} (−0.270). Although, the total diversity ($H = 1.5$) is lower than the weed community in a farmland ($H = 2.87$) [29]. \textit{M. longibracteatus} is more prominent in sandy mangrove shore as revealed in previous study [30].

### 3.5. Plant Species Importance Value

The result (Table 5) indicate that \textit{Mariscus longibracteatus} (Family: cyperaceae) are the most abundant ($n = 1570$), and thus have the highest importance value in the study site ($I_v = 58.24$). The order of importance of weed species found in

### Table 3. Mean levels of total hydrocarbon content (THC) and heavy metals ± 1 SE in weed collected Eagle Island, Niger Delta Nigeria.

| Species        | Cadmium  | Lead    | Zinc     | THC     |
|----------------|----------|---------|----------|---------|
| \textit{A. areum} | 0.002 ± 0.0005 | 3.01 ± 0.12 | 52.67 ± 0.45 | 21.09 ± 0.23 |
| \textit{C. longipinna} | 0.002 ± 0.0005 | 3.01 ± 0.13 | 49.96 ± 0.07 | 20.16 ± 0.38 |
| \textit{C. mindorensis} | 0.23 ± 0.005 | 3.36 ± 0.01 | 51.31 ± 1.31 | 21.55 ± 1.05 |
| \textit{D. oliveri} | 0.002 ± 0.0005 | 3.05 ± 0.15 | 49.55 ± 0.69 | 11.61 ± 0.94 |
| \textit{M. ligularis} | 0.26 ± 0.005 | 1.87 ± 0.01 | 13.44 ± 0.01 | 21.85 ± 0.02 |
| \textit{M. longibracteatus} | 0.002 ± 0.0005 | 3.11 ± 0.11 | 51.72 ± 0.11 | 6.30 ± 0.02 |
| \textit{P. vaginatum} | 0.14 ± 0.005 | 5.53 ± 0.01 | 19.82 ± 0.01 | 2.91 ± 0.40 |
| \textit{R. indica} | 0.24 ± 0.005 | 5.97 ± 0.01 | 30.77 ± 0.01 | 3.03 ± 0.03 |
| \textit{S. indica} | 0.001 ± 0.00 | 3.45 ± 0.15 | 51.18 ± 0.15 | 15.21 ± 0.33 |
### Table 4. Diversity of plant species in sand fill mangrove forest at Eagle Island, Nigeria.

| Scientific name | Common name                | Abundance | $p_i$ | $\ln(p_i)$ | $p_i \times \ln(p_i)$ |
|-----------------|----------------------------|-----------|-------|------------|------------------------|
| *Mariscus ligularis* | NA                        | 900       | 0.232 | −1.461     | −0.339                 |
| *Paspalum vaginatum* | Seashore paspalum          | 525       | 0.135 | −2.002     | −0.270                 |
| *Reissantia indica* | NA                        | 505       | 0.130 | −2.040     | −0.265                 |
| *Cyperus mindorensis* | White water sedge         | 50        | 0.013 | −4.343     | −0.057                 |
| *Mariscus longibracteatus* | NA                       | 1570      | 0.405 | −0.904     | −0.366                 |
| *Acrostichum aureum* | Mangrove fern              | 212       | 0.955 | −0.046     | −0.044                 |
| *Calamus longipinna* | NA                        | 99        | 0.026 | −3.650     | −0.095                 |
| *Daniellia oliveri* | Ilorin balsam              | 5         | 0.001 | −6.908     | −0.007                 |
| *Stachytarpheta indica* | Blue snake weed           | 10        | 0.003 | −5.809     | −0.017                 |
| *Solanum torvum Swartz* | Prickly solanum         | 3         | 0.001 | −6.908     | −0.007                 |
| *Emilia praetermissa* | Yellow tassel flower       | 5         | 0.001 | −6.908     | −0.007                 |
| *Luffa cylindrica* | Loofah                     | 7         | 0.002 | −6.215     | −0.012                 |
| *Oldenlandia corymbosa Linn.* | NA                     | 3         | 0.001 | −6.908     | −0.007                 |
| *Corchorus olitorius L.* | Nalta jute              | 2         | 0.001 | −6.908     | −0.007                 |
| *Terminalis catappa* | Indian almond              | 4         | 0.001 | −6.908     | −0.007                 |
| **SUM**          |                           | 3880      | 1.00  |             |                        |

$H = −1.507$

### Table 5. Abundance, importance value ($I_v$), relative density, relative frequency and relative dominance for Eagle Island in the Niger River Delta, Nigeria.

| Scientific name | Abundance | Relative density (%) | Relative frequency (%) | Relative dominance $I_v$ |
|-----------------|-----------|----------------------|------------------------|--------------------------|
| *Mariscus ligularis* | 900       | 23.20                | 15.56                  | 1.14 × 10⁻⁵              | 38.75                   |
| *Paspalum vaginatum* | 525       | 23.20                | 13.33                  | 4.15 × 10⁻⁶              | 36.53                   |
| *Reissantia indica* | 505       | 23.20                | 11.11                  | 5.55 × 10⁻⁵              | 34.31                   |
| *Cyperus mindorensis* | 50        | 1.29                 | 6.67                   | 2.93 × 10⁻⁵              | 7.96                    |
| *Mariscus longibracteatus* | 1570     | 40.46                | 17.78                  | 1.65 × 10⁻⁵              | 58.24                   |
| *Acrostichum aureum* | 212       | 5.46                 | 8.89                   | 1.03 × 10⁻⁴              | 14.35                   |
| *Calamus longipinna* | 99        | 2.55                 | 6.67                   | 2.25 × 10⁻⁵              | 9.22                    |
| *Daniellia oliveri* | 5         | 0.13                 | 2.22                   | 1.83 × 10⁻⁴              | 2.35                    |
| *Stachytarpheta indica* | 10        | 0.26                 | 4.44                   | 4.59 × 10⁻⁵              | 4.70                    |
| *Solanum torvum Swartz* | 3         | 0.08                 | 2.22                   | 3.71 × 10⁻⁵              | 2.30                    |
| *Emilia praetermissa* | 5         | 0.13                 | 2.22                   | 1.14 × 10⁻⁴              | 2.35                    |
| *Luffa cylindrica* | 7         | 0.18                 | 2.22                   | 5.02 × 10⁻⁴              | 2.40                    |
| *Oldenlandia corymbosa Linn.* | 3         | 0.08                 | 2.22                   | 1.65 × 10⁻⁵              | 2.30                    |
| *Corchorus olitorius L.* | 2         | 0.05                 | 2.22                   | 7.36 × 10⁻⁴              | 2.27                    |
| *Terminalis catappa* | 4         | 0.10                 | 2.22                   | 5.62 × 10⁻⁴              | 2.33                    |
| **SUM**          | 3880      |                      |                       |                          |                         |
sand fill soil include; *M. longibracteatus* > *M. ligularis* > *P. vaginatum* > *R. indica*. All species are weed except *Acrostichum aureum* L., which is classified as a mangrove species under the family of Pteridaceae [31]. They are however, regarded as weed in some places, for instance in Matang, Malaysia [32]. *P. vaginatum* are also regarded as invasive species [33].

4. Discussion

Mangroves are habitat specific, and grow only in swampy soils. This is because most mangrove species apart from *A. aureum* (mangrove fern) cannot grow well in sandy soil because it has low salinity and conductivity (Table 1). Mangrove swamp is one of the largest carbon sequesters in the world [34] [35], this is because of their air purification ability and high productive capability [17]. Swampy soils have higher heavy metal load because of their exposure to oil spillages from oiling activities onshore and offshore. Pollution of the shorelines destroys swampy soils by reducing salinity and destroying microbes within the soil. A known characteristic of mangrove swamp is their ability to carry out decomposition [36], which has made them a biodiversity hot spot [37]. But when human activities of deforestation, sand mining and urbanization degrade the soil it becomes difficult for them to carry out their function as host to numerous soil dwelling organisms. This enables opportunistic invasive species and weeds to come in to colonize the area [38]. In this study, previous sand mining activity destroyed the swampy soil, and in its place sandy soil was deposited, which facilitated the proliferation of weeds [30]. Growth of weeds near or inside mangrove forest is becoming a hallmark of a disturbed forest [27]. The role played by weeds in this study area is not well known but need further studies. However, the result indicates that weeds present in the sand filled area absorbed the soil pollutants. This is shown by Figure 3, where weed concentration of THC and heavy metal was more than that of the soil. It shows that the weeds are acting as bioremediation agents. Mangrove swamp has hydrocarbon utilizing bacteria that degrade pollutant to a less harmful level [39] [40]. The weed species from field observations are acting as host to pollinators such as butterflies, cricket, beetles and mosquitoes, to mangrove; this is a positive role by weeds [41]. However, they can be detrimental to humans by hosting harmful insects and rodents. Field experiences had shown that most mosquitoes that reside in mangrove forest always go to roost on nearby weed and enter the mangroves to feed on animals, which is a negative role because of the proliferation of mangroves parasites in Africa [42].

Most of the weeds found in this study are dominant because they are all aquatic and lowland weeds that benefit from the closeness of the area to a river. The change in the habitat from swampy to sandy soil contributed to the transition from exclusively a mangrove community to a weed community. Loss of mangrove forest as a result of anthropogenic action lead to a loss of the ecosystem services they provide e.g. carbon sequestration, fire wood and basket production.
and fisheries. The weed are not known to provide any significant ecosystem services to the environment apart from being host to some disease parasites, causing bush fires and acting as a nuisance in the environment. The species *Mariscus longibracteatus* are more dominant because they are aquatic weed that grow in wet grounds in forest zones. They are of the family Cyperaceae [2]. They are large tufted sedge that can grow to about 1 m, and are produced from seeds. Their seeds are tiny and are propagated by wind. They have very prominent spikes on an inflorescence, which sticks to clothing and facilitates dispersal and propagation. This has made them to be found in any environment. Although they are mangrove forest plants, but are introduced through human activities such as logging, sand filling, construction and reclamation. Their presence in a mangrove forest shows that the system had been disturbed by humans.

High concentration of Zn can be ascribed to increased land runoffs and influx of metal-rich water in the weed root soil giving rise to elevated metal levels. Similarly, elevated levels of metals in weed root soil were observed in Pondicherry Harbor [43]. However, it was found that the total concentrations of all selected heavy metals in the weed root soil were below the critical soil concentration values [44], which reflect the topography and bed rock of the area as the origin of these metals.

5. Conclusion and Recommendation

This study is significant because it shows that human activities are major causes of weed invasion of mangrove forest, and weeds are primary successors in disturbed environment. Change in soil condition can lead to the elimination of mangroves due to their affinity for swampy soil rather than sandy soil. To revert the invasion of mangrove forest by foreign weed species, we need to create a protective barrier around the forest to prevent the encroachment of humans. As a way of restoring the mangrove forest there can be a replacement of the sandy soil with swampy soil. Furthermore, weeds can be used to monitor THC and heavy metal contamination of mangrove forest.

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Conflicts of Interest

The author declares no conflicts of interest regarding the publication of this paper.

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