Floating treatment wetlands for domestic wastewater treatment

ABSTRACT
Floating islands are a form of treatment wetland characterized by a mat of synthetic matrix at the water surface into which macrophytes can be planted and through which water passes. We evaluated two matrix materials for treating domestic wastewater, recycled plastic and recycled carpet fibers, for chemical oxygen demand (COD) and nitrogen removal. These materials were compared to pea gravel or open water (control). Experiments were conducted in laboratory scale columns fed with synthetic wastewater containing COD, organic and inorganic nitrogen, and mineral salts. Columns were unplanted, naturally inoculated, and operated in batch mode with continuous recirculation and aeration. COD was efficiently removed in all systems examined (> 90% removal). Ammonia was efficiently removed by nitrification. Removal of total dissolved N was ∼50% by day 28, by which time most remaining nitrogen was present as NO3-N. Complete removal of NO3-N by denitrification was accomplished by dosing columns with molasses. Microbial communities of interest were visualized with denaturing gradient gel electrophoresis (DGGE) by targeting specific functional genes. Shifts in the denitrifying community were observed post-molasses addition, when nitrate levels decreased. The conditioning time for reliable nitrification was determined to be approximately three months. These results suggest that floating treatment wetlands are a viable alternative for domestic wastewater treatment.

Key words | bacteria, biofilm, COD, denitrification, floating treatment wetland, nitrification

INTRODUCTION
A type of artificial wetland in which emergent plants are grown either hydroponically or in a media floating on the surface of a pond-like basin has been used for habitat enhancement or contaminant amelioration since at least the mid-1970s (Seidel & Happel 1986; Hoeger 1988). As with more conventional surface flow and subsurface flow treatment wetlands, Floating Treatment Wetlands (FTW) have been employed for removal and treatment of a diverse array of contaminants and polluted waters (Headley & Tanner 2008b). However, due to their ability to float with relatively large fluctuations in water level, treatment of stormwater (Headley & Tanner 2008a, b) and combined sewer overflow (Van de Moortel et al. 2010b) appear to be the most typical applications. Natural floating islands, which can exist in locations where plant growth exceeds decay rates, maintain buoyancy via trapped gasses within a matrix of partially decayed and living plant material (Mitsch & Gosselink 2000). Ayaz & Saygin (1996) describe a completely hydroponic system, but most other FTW employ a superstructure frame constructed of buoyant material (PVC, bamboo, polystyrene, etc.) (Hoeger 1988; Billore et al. 2008; Van de Moortel et al. 2000a, 2010b), are supported by cables attached to the bank (Kerr- Upal et al. 2000), or are constructed from an inherently buoyant planting media (Todd et al. 2005; Headley & Tanner 2008a; Stewart et al. 2008). Depending on the thickness of the planting media and species of emergent plants employed, roots can be completely contained within the media or extend through and be exposed directly to the water column below.

Compared to other treatment wetland systems, design of FTW is based on very limited information and most applications seem to be unique for even the most basic parameters such as size, degree of buoyancy, planting media, plant selection, etc. Only a few studies (e.g. Headley &
Tanner 2008a; Nakai et al. 2008; Li et al. 2010; Van de Moortel et al. 2010a, 2010b) have attempted to assess performance in replicated experiments and, due to the variety of designs and wastewater types and treatment objectives, performance generalizations are not possible at this time. To further advance understanding of processes important in FTW for domestic wastewater applications, we have been conducting experiments on chemical oxygen demand (COD) and nitrogen removal and associated microbial populations in FTW. An important criterion in FTW design is the type of planting media (matrix). An inherently buoyant matrix avoids the use of a supporting frame and we have focused attention on various buoyant matrix materials available through a local commercial provider. This paper summarizes research on the removal of COD and nitrogen species together with the dynamics of microbial communities occurring within two different matrix materials. The non-woven matrix is made of 100% recycled plastic fabricated into floating mats which can be configured, as shown in Figure 1, to include pumps for recirculation as well as aeration systems at various points within the matrix. By varying flow rate, duration and frequency of recirculation and aeration it is possible to control nutrient loading rates and redox conditions within the FTW matrix.

Previous research with these matrix materials conducted in outdoor ponds provided observations of the substantial disappearance of key wastewater constituents including COD, ammonia, nitrate and phosphate (Stewart et al. 2008). Since plants were not incorporated, constituent removal in these outdoor experiments was likely due to the activity of microorganisms growing as biofilms on surfaces within the island matrix. The objective of the current study is twofold: (1) determine the optimum operational conditions to encourage simultaneous nitrification (ammonia removal) and denitrification (nitrate removal) within a FTW environment by stimulating the appropriate microbial communities; and (2) determine the microbial community response to variations in aeration, matrix material and organic carbon loading. Successful completion of this research will not only provide the basis for improving FTW design and efficacy, but will provide insight into the processes responsible for effective water quality remediation occurring within FTW.

**METHODS**

Experiments were conducted in laboratory scale systems consisting of 20 cm diameter columns containing matrix material 20 cm thick submerged 10 cm below the water surface. Matrix material was either a very porous commercial mat made from 100% recycled plastic or loose shredded carpet fibers contained within a porous mesh, both supplied by Floating Island International (www.floatingislandinternational.com). Additionally, two otherwise identical columns, one filled with 20 cm pea gravel, the other left as open water, were included for comparison. Columns were each filled with 20 liters of simulated domestic wastewater containing ~500 mg/L COD (mostly from sucrose), ~15 mg/L NH₄-N and ~15 mg/L NO₃-N, ~30 mg organic N/L from Primatone (Sigma), and other inorganic components (Taylor et al. 2011). All columns were unplanted, inoculated with soil and pond water, and operated in batch mode with continuous recirculation from the bottom of the column (at 20 mL/min) with continuous aeration (unless noted) from aquarium pumps into the surface water 10 cm above the matrix. We summarize data from five consecutive batch runs (B1–B5) that followed a conditioning period of four batches over a three month period. B1, B2, and B3 were run for 28 days. B4 and B5 were run for 42 days. During B4, columns were dosed with 10 g molasses (measured as 820 mg COD/g) on day 25 and again on day 29 in order to provide reducing equivalents for denitrification. There was no aeration during days 29–42.
Water quality analysis

Water samples were collected from the recirculating return flow on days 0, 3, 8, 14, 21, and 28 of batches B1–B3. Additional sampling during B4 and B5 was on days 35 and 42. Samples were filtered through a 0.2 μm PES filter and stored in glass scintillation vials (4°C). Samples were analyzed for COD, NH₄-N, and total N using HACH (Loveland, CO) methods. NO₂-N and NO₃-N were measured by ion chromatography (Dionex Corp., Sunnyvale, CA).

Floating Island biofilm collection and DNA extraction

Biofilm samples were collected on Day 0 of B3 (by which time columns were well conditioned) and again on Day 0 of B5 (14 days after the second molasses dose during B4) from three depths within each treatment: top (upper 5 cm of material), center (middle 5 cm of material), and bottom (lower 5 cm of material). To clearly distinguish between samples collected on Day 0 of B3 (by which time columns were well conditioned) and again on Day 0 of B5 (14 days after the second molasses dose during B4) from three depths within each treatment: top (upper 5 cm of material), center (middle 5 cm of material), and bottom (lower 5 cm of material). To clearly distinguish between depths sampled, a 2.5 cm zone was left unchanged between each of the locations. The open water column was sampled by vacuum filtering 250 mL of effluent through a 0.2 μm polycarbonate membrane. Field samples from unplanted FTW, operated by Floating Island International, were also provided and analyzed for comparison with our laboratory samples. Materials collected from each treatment were placed directly into MO BIO PowerBead Tubes (MO BIO PowerSoil™ DNA Isolation Kit). The PowerSoil™ DNA Isolation Kit was used to complete the DNA extraction as described in the manufacturer’s protocol with the exception that PowerBead tubes were placed into the FastPrep® Instrument (Qbiogene, Inc.) at speed 5.5 for 45 s. DNA yield was estimated on an agarose gel with ethidium bromide and was used for DGGE. Additional sampling during B4 and B5 was on days 35 and 42. Samples were paired with primer RottF for amplifying fragments to be analyzed by denaturing gradient gel electrophoresis (DGGE). Presumptive presence of the amoA gene was indicated on an agarose gel by a 531 bp PCR product. PCR reactions (20 μL) were performed using 2X GoTaq® Green Master Mix (www.promega.com). The PCR reaction mixture consisted of 10 μL 2X GoTaq® Green Master Mix, 0.5 μL Ultrapure BSA (50 mg/mL, Ambion), 2.5 μL DEPC-treated water, 1 μL 12.5 μM forward and reverse primer, and 5 μL 1:10 diluted (unquantified) template DNA. PCR amplifications were performed on an Eppendorf Mastercycler® ep thermal cycler (Eppendorf North America, www.eppendorfna.com) using the following program. An initial denaturation for 60 s at 94 °C was followed by a total of 35 cycles of amplification consisting of denaturation at 94 °C for 60 s, annealing at 54 °C for 60 s, and extension at 72 °C for 3 min. The program ended with an extension step at 72 °C for 10 min (Bahr et al. 2005). PCR products were confirmed by agarose gel electrophoresis and staining with ethidium bromide and were used for DGGE.

PCR targeting functional genes

Ammonia monoxygenase gene

Oligonucleotide primers were synthesized by Integrated DNA Technologies (www.idtdna.com). PCR primers NirS cd3aF (5’ GTSAACTSAAGGARACSGG 3’, Michotey et al. 2000) and NirS R5cdR (5’ GASTTGGRTGSGTC TTGA 3’, Throback et al. 2004) along with NirK F1aCuF (5’ ATCATGTTCTGCCGCGC 3’, Hallin & Lindgren 1999) and NirK R3CuR (5’ GGCCTGATCAGRTTGGTT 3’, Hallin & Lindgren 1999) target the two forms of the nitrite reductase gene (nir, required for nitrite reduction to nitric oxide). Primers NirS R3cdR and NirK R3CuR were synthesized with a 5’ 40-bp GC clamp (described above) and were paired with their respective forward primers for amplifying fragments to be analyzed by DGGE. Presumptive presence of the nirS and nirK genes was indicated on an agarose gel by a 465 bp or a 502 bp PCR product, respectively. PCR amplifications were performed on an Eppendorf Mastercycler® ep thermal cycler (Eppendorf North America, www.eppendorfna.com) using the following program. An initial denaturation for 2 min at 94 °C was followed by a total of 35 cycles of amplification consisting of denaturation at 94 °C for 30 s, annealing at 57 °C for 60 s, and extension at 72 °C for 60 s. The program ended with an extension step at 72 °C for 10 min (Throback et al. 2004). PCR products were confirmed by agarose gel electrophoresis and staining with ethidium bromide and were used for DGGE.
Denaturing gradient gel electrophoresis

DGGE was performed on PCR products from community DNA using a DCode™ system (www.biorad.com) and reagents from Sigma-Aldrich (www.sigmaaldrich.com). Gels had a gradient of denaturant concentrations from 40% at the top of the gel to 70% at the bottom, where 100% denaturant is defined as 7 M urea and 40% formamide. Gels also contained an 8–12% polyacrylamide gradient from top to bottom (Girvan et al. 2005). Electrophoresis was at 60 V for 16 h. Gels were stained with Sybr®Gold (www.invitrogen.com) and documented using a FluorChem™ 8800 fluorescence imager (www.alphainnotech.com). Three marker lanes (generated from five pooled clones) were included in each DGGE gel so that lane profiles within and between gels could be compared. Bands in DGGE images were identified visually on a presence–absence basis. Band intensities were not physically measured, but visually prominent bands were considered to represent numerically significant members of the community.

DGGE data analysis

DGGE gels were compared and analyzed for each gene investigated using the GelCompar II software (Version 6.1, Applied Maths Inc.). Subsequent statistical analyses were performed using R software libraries labdsv (Roberts 2009) and optpart (Roberts 2010) (www.r-project.org). Similarity and dissimilarity matrices were calculated using Dice (Gel-Compar II) and Sorensen (R). Hierarchical clusters were generated from these matrices and displayed using the Unweighted Pair Group Method using Arithmetic averages (UPGMA). This method displays the average similarity between profiles for each sample compared. Hierarchical clusters displayed were generated from the similarity matrix calculated using the GelCompar II software; while more robust statistical analyses were performed using the dissimilarity matrix calculated using the R software. Reported large D² values (similar to R²) indicate more confidence in the resultant clusters whereas small p-values (determined using the chi-squared test) indicated that the clusters represented the dataset well.

RESULTS AND DISCUSSION

Water quality

Because all water quality analyses were done on filtered samples (0.2 μm pore size), bacterial cells were excluded. The laboratory columns containing plastic matrix, carpet fibers, pea gravel or open water were all effective at removing COD and nitrogen (Figure 2). There was relatively little difference among treatments (except as noted below). COD removal in all treatments was ~90% within the first two weeks of each batch, i.e., from ~500 mg COD/L initially to <50 mg COD/L by day 14. Initial total dissolved nitrogen was ~60 mg N/L, consisting of ~30 mg N/L organic N (from Primatone), ~15 mg NO3-N/L, and ~15 mg NH4-N/L. Total dissolved N generally decreased by ~50% within the first two weeks, but leveled off after that. In the first week, removal of NO3-N was usually ~90%, probably from denitrification. NH4-N usually increased in the first week, probably from mineralization of organic N. An exception was the gravel column where NH4-N decreased immediately. This

![Figure 2](https://iwaponline.com/wst/article-pdf/64/10/2089/444032/2089.pdf)
behavior may have been the result of adsorption onto the gravel matrix and has been observed in previous studies using the same gravel (Riley et al. 2005). By days 21–28, NO₃-N usually began to accumulate again and often accounted for most of the total N. There was usually a corresponding loss of NH₄-N during this time period, indicating that nitrification was occurring. Differences in nitrification between new and conditioned plastic matrix suggested that about three months were required to establish an effective nitrifying biofilm community. By day 28 in batches B1-B3, the columns had reached a steady state in which COD was virtually absent, and almost all of the total dissolved N was as NO₃-N (~20 to 30 mg N/L). To test the hypothesis that denitrification in these batches had been carbon-limited, we introduced doses of molasses (10 g molasses/column) on days 25 and 29 of B4. Dosing produced a spike in COD (to ~370 mg COD/L), but by the end of the batch two weeks later, >90% of this COD had also been removed. The molasses was effective at increasing denitrification. By the end of the batch on day 42, total dissolved N was <5 mg N/L, NO₃-N was <4 mg N/L, and NH₄-N was <2 mg N/L. COD was <100 mg/L, but this residual COD from the molasses would probably have been removed had the batch been allowed to run beyond day 42. Batch B5 produced results that were similar to B4. NO₂-N was not a component of the synthetic wastewater and was usually ≤1 mg N/L for all treatments and time points.

Synthetic wastewater (completely soluble) was used in these column experiments. Additional pretreatment may be necessary when implementing FTW for wastewaters containing grit and suspended solids, etc.

Microbial community responses

The responses of the nitrifying and denitrifying microbial communities were monitored using DGGE. Interpretation of DGGE profiles was done cautiously as they are invariably a mix of artifact and real diversity. Individual bands are generally assumed to represent individual genotypes, but only DNA sequencing can confirm this. The total number of bands in a profile is a rough estimate of diversity and the intensity of a band is a rough estimate of the prominence of the corresponding genotype in the microbial community (Muyzer et al. 1993). It should be noted that bands suspected to be artifact were not included in the analysis and as a result, diversity may have been underestimated. Analysis of DGGE gels was performed using Gel Compar II software (v. 6.1, Applied Maths Inc.) to visualize and compare gels.

Statistical analyses were performed based on band presence/absence within each profile using the R software libraries labdsv (Roberts 2009) and optpart (Roberts 2010) (www.r-project.org).

Nitrifying community

The nitrifying communities within each column were observed to have limited diversity (maximum of 12 bands observed, gravel treatment, Figure 3). The open water column had developed a distinct nitrifying community unlike any of the other treatment conditions ($D^2 = 0.9944$). Dosing with molasses and ending aeration did not significantly affect the structure of the nitrifying communities. The community structure of the laboratory versus field samples were distinctly different ($D^2 = 0.9955$). Field samples generally contained 5–7 bands, while laboratory samples contained 5–12 bands. It appeared that sample depth within the column was the most significant factor in community structure ($p < 0.001$, Figure 3) followed by FTW material used ($p = 0.01$), with matrix and carpet communities being more similar to one another than to the gravel community.

Denitrifying community

In order to investigate the entire denitrifying community present, both the nirS and nirK genes were characterized. Overall, the denitrifying community profiles were
considerably more diverse compared to the nitrifying community profiles. As observed with the nitrifying community profile, the profiles for the open water column had developed unique communities compared to the other treatments ($D^2 = 1$, for both genes). For the nirK gene, the communities were highly diverse, but apparent similarities were specific to the FTW matrix material within the column ($D^2 = 0.9996$, $p = 0.005$). For the nirS gene, the FTW matrix material was most important in determining the community that developed ($p < 0.001$) with all of the gravel samples grouping onto a single branch. Adding molasses and ending aeration also appeared to affect the nirS denitrifying community, though not as greatly as FTW material had ($p = 0.05$, Figure 4). Finally, the field samples had developed significantly different denitrifying communities for the nirS ($p = 0.001$) but not the nirK gene.

FTW matrix material had the largest effect on the denitrifying community. As observed with the nitrifiers, the elimination of aeration and addition of molasses did not significantly affect the community structure, but did stimulate denitrifying activity and thus nitrate removal.

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