Fiber Attachment Module Experiment (FAME): Using a Multiplexed Miniature Hollow Fiber Membrane Bioreactor Solution for Rapid Process Testing

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Introduction:
- Bioreactor research is mostly limited to continuous stirred-tank reactors (CSTRs) which are not an option for microgravity (µg) applications due to the lack of a gravity gradient to drive aeration as described by the Archimedes principle. Bioreactors and filtration systems for treating wastewater in µg could avoid the need for harsh pretreatment chemicals and improve overall water recovery.
- Solution: Membrane Aerated Bioreactors (MABR) for µg applications, including possible use for wastewater treatment systems for the International Space Station (ISS).
- Small 1-4 volume MABRs do not lend themselves to rapid testing capacities. To address this, we designed 125-ml, rectangular reactors, to create the Fiber Attachment Module Experiment (FAME) system.
  - Each FAME module (i.e., reactor) is self-contained.
  - Easy to use with pumps for continuous recycling of fluids/feeding and sensors for parameter monitoring similar to their larger counterparts.
- Application: Rapid Biofilm Attachment Studies
  - Goal: Achieve rapid reactor startup and biofilm attachment (based on carbon oxidation and nitrification of wastewater).
  - Multi-factorial test with data collected over short to study and improve bioreactor performance.
  - Thirteen fiber surface treatments and three inoculation sources tested.

Biological Results:
- TTU inoculum showed significantly higher attached cell counts compared to R3 inoculum in all treatments except HF - 15 min.
- Highest cell counts overall were seen for TTU inoculum on FE – 18 sec & HF – 12 sec fibers.
- R3 counts were the lowest; no significant difference between TTU and Septic.
- R3 inoculum was effluent from well-established bioreactor; selecting for planktonic cells in the effluent could explain lower attachment.
- Septic tank inoculum was compared to test a local and consistent source for future studies if comparable to TTU inoculum results.

Materials and Methods:
- Reactor Setup:
  - Multiple polycarbonate FAME racks, each containing 4 modules
  - Peristaltic pumps used for fluid recirculation (1025 reactor volumes per day)
  - Standard breathing air used as reactor oxygen source
  - Modules contained in a Controlled Environment Chamber (CEC) at 25°C in darkness

Fiber Treatments:

<table>
<thead>
<tr>
<th>Fiber Treatments</th>
<th>Etching</th>
<th>Polymer Coating</th>
<th>Mechanical</th>
<th>Scaffolding</th>
</tr>
</thead>
<tbody>
<tr>
<td>3% HF Fluoroetch®</td>
<td>Polystyrene Acid (PAA)</td>
<td>Abrasion via Sand Paper</td>
<td>Nylon Cable</td>
<td>5K Polyacrylic Acid (PAA)</td>
</tr>
<tr>
<td>Abrasion via Silk</td>
<td>Polyester (PS)</td>
<td>Sheaths</td>
<td>Silk Thread</td>
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<td>FE Silk</td>
<td>Spider Silk</td>
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Inoculation Sources:
- Effluent from a KSC single-stage carbon oxidation reactor (R3)
- Effluent from a Texas Tech University (TTU) single-stage MABR
- Activated sewage sludge from a local septic tank

Evaluation Parameters:
- Bulk fluid analysis weekly: NH4, NO3, NO2, total cell counts
- Sample volumes replaced with reactor feed (real urine solution)
- End of experiment: total cell counts for biofilms attached to fibers
- Oxygen permeation of modified fibers compared to controls

Chemical Analysis Results:
- Limited chemical analysis performed due to nature of experiment; bulk fluid samples collected weekly for pH, NH4, NO3, and NO2; pH for reactor inocula and feed ranged between 6.3 and 6.5; as experiments progressed, pH rose rapidly to between 8 and 9 due to urea hydrolysis to ammonia.
- Buildup of NH4 indicative of no nitrification; supported by absence of nitrate and nitrite species. Attempt made to lower pH to induce nitrification at DOE 40; attempts were unsuccessful.

Results show that formed biofilms were heterotrophs responsible for carbon oxidation/urea hydrolysis. Further fiber development may be required to better attract nitrifying biofilms.

Oxygen Permeation Results:
- PDMS fibers traditionally used for their high O2 permeation, superior mechanical/chemical resistance, and toleration of high intra-membrane pressures.
- Modification of PDMS for better biofilm attachment must not sacrifice these properties, especially O2 permeation.
- O2 permeation comparisons completed for five of the modification processes:
  - Mass transfer coefficient, K, derived for each membrane type.
  - Describes resistance between gas phase, membrane, and liquid phase boundary layers.
  - Increased K-quantities to increased O2 transfer across the membrane.
- None of the modifications exhibited statistically different K values, showing O2 permeation was not inhibited by the treatment.
- Further shown that differences in biofilm attachment are not due to changes in O2 permeation.

Future Work:
- Future experiments involving FAME hardware will include further testing of fiber modifications, inoculum choice, reactor feed composition, reactor poisoning effects, and oxygen permeation treatments, and more.
- Fiber modifications which have shown promise will also undergo further verification testing.
- Modifications to current hardware include the addition of external probe tanks for continuous monitoring of pH and dissolved oxygen (DO) within the modules to allow for improved process controls in hopes of gaining nitrification in the reactors.
- The value of this hardware is demonstrated for rapid testing of numerous parameters in parallel. FAME racks may also be utilized for future reactor hibernation and biofilm-development time course studies.