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**Recent Work** 

## Title

Application of custom-designed fermentors for extremophilic microorganisms

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VIMSS Virtual Institute for Microbial Stress and Survival

# Abstract

Background: Extremophilic microorganisms may play key roles in remediation of groundwater contaminants and biofuels development Standard fermentors are not equipped for anaerobic growth conditions, nor can stainless steel withstand the reactivity of metal-reducing organisms and their metabolic by-products. We have developed methods to grow the sulfate-reducing bacterium, Desulfovibrio vulgaris Hildenborough (DvH), under various, batch and continuous growth conditions for detailed physiological and molecular analyses, using custom-built fermentors.

Methods: Five-liter-volume fermentors have PEEK headplates and agitators. During growth, temperature, pH, OD, and redox potential are continuously controlled. Samples are taken for ion chromatography, phospholipid fatty acid (PLFA) analysis, direct cell counts, RT-PCR, and total cell protein. DvH is grown in batch and turbidostat modes using a defined, lactate-sulfate medium at 30°C. Batch cultures have also been grown under various stress conditions.

**Results:** The maximum specific growth rate for DvH in the 5-liter fermentor is 0.11 h<sup>-1</sup>. At the onset of deceleration phase, all lactate (60) mM) is depleted, and 30 mM of the initial 50 mM sulfate is metabolized. The dilution rate to maintain the culture at mid-log phase in the turbidostat is 0.15 h<sup>-1</sup>. Cell densities and total proteins range from 5-10 x 10<sup>8</sup> cells/ml, and 80-120 µg/ml, respectively. PLFA profiles for DvH are sensitive to the growth conditions and growth phase, while preliminary results show consistent whole-cell protein patterns when visualized using SDS-PAGE. Melt curves generated by real-time PCR proved to be an excellent quality control tool to monitor culture purity. **Conclusion:** Detailed characterization of DvH during growth in custom-designed pilot-scale fermentors provides insights into physiological changes during different growth phases and stress conditions, and has resulted in the development of protocols for controlled and reproducible production of high quality biomass. The experiments have also proven the utility of custom-designed fermentors for growth of extremophiles.

# **Description of Fermentors**





Two fermentors were run simultaneously in turbidostat mode to produce 100-liter DvH/week for the Protein Analysis Complex Project. The 10-liter medium bottle is shown between the fermentors. Effluent was continuously collected and chilled to 4°C (right image), while cells were harvested by batch centrifugation. Anaerobic headspace is maintained by flow of sterile nitrogen, or other anaerobic gas mixes.

# **Batch Growth - Effect of initial lactate concentration**

Growth of DvH in lactate/sulfate medium with 10% inoculum was compared for initial lactate concentrations of 60, 80 and 100 mM. The initial sulfate concentration was 50 mM in all cases. pH was maintained at 7.2 with 1 N HCI. The mixing rate was set to 200 RPM. Redox potential increased from -700 to -450 mV during growth phase, and furthermore to -350 mV during stationary phase.





Higher protein concentrations were measured for 100 mM initial lactate

hours since inoculation

# Application of custom-designed fermentors for extremophilic microorganisms

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universal primer set, melt

temperature: 84.5°C

DvH-specific primer set,

melt temperature: 86.5°C

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