SUNLIGHT SUPPLY AND GAS EXCHANGE SYSTEMS IN MICROALGAL BIOREACTOR

K. Mori*, H. Ohy?, K. Matsumoto**, and H. Furune***

*Keio University, 3-14-1 Hiyoshi, Kohoku-ku, Yokohama 223, Japan; **Yokohama National University, Tokiwadai, Hodogaya-ku, Yokohama 240, Japan; ***La Foret Engineering and Information Service Co., 2-7-8 Toranomon, Minato-ku, Tokyo 105, Japan

ABSTRACT

The bioreactor with sunlight supply system and gas exchange systems presented here has proved feasible in ground tests and shows much promise for space use as a CELSS device.

Our chief conclusions concerning the specification of total system needed for a life support system for a man in a space station are the following.

1. Sunlight supply system : compactness and low electrical consumption.
2. Bioreactor system : high density and growth rate of chlorella.

INTRODUCTION

The sunlight supply system /1-2/ presented will provide the visible solar radiation necessary for the various activities in the space station, such as cultivation experiments on algae, plants, fishes, birds, and animals room, lighting for modules, and crew sun-bathing. Even natural solar rays reaching the earth’s surface contain ultraviolet rays of medium and long wavelengths (UV-B and UV-A) and infrared and heat rays, all of which are harmful to life. On a space station, the most dangerous short-wavelength ultraviolet rays (UV-C), x-rays, and gamma-rays are added, and those cited above are present in markedly higher intensity. It will thus only be after elimination of these harmful rays that special solar rays will be utilisable in the space station for activities related to the life sciences. With respect to the Regenerative Life Support System in CELSS, the main function of the microalgal bioreactor with gas exchange system to supply O2 and assimilate exhaust of CO2 for a man in a space station, though another would be to produce algal biomass for a human food and medicine: the biomass protein, carbohydrate, and β-carotene would be useful.

CONCEPT AND BASIC DESIGN OF TOTAL SYSTEM

Sunlight Supply System

The basic design of the sunlight supply system is shown in Figure 1 /1/. The solar rays are collected and concentrated by a factor of 10,000 by means of a solar collector assembly composed of many small Fresnel lenses.

Limitation of the admitted spectral band to the visible region (380-770 nm wavelength) is made possible by the chromatic aberration of the individual Fresnel lenses. Chromatic aberration caused by the Fresnel lenses separates the focal points of different wavelengths, so that placing the end of the fiber optic cable at the focus of green light admits mainly the visible spectrum (Figure 2).

The solar rays focused by each small lens are fed through the well-polished terminal end of a flexible light-conducting fiber, and are confined within its quartz core for transmission to the required location.

Solar collection is performed by Fresnel lenses assembled in a honeycomb lattice as illustrated in Figure 1. The front face of each is covered by a protector serving also as a filter. The protector is made of quartz, is resistant to scratching, and is sufficiently thick to withstand most impinging meteoroids, debris and deposits that may be encountered. Moreover, in the event of damage to the protector, the affected unit can be cut off and sealed off from the rest of the system, which can continue functioning independently. The space between protector and array
independently. The space between protector and array is filled with an inert gas to allow temperature control and to suppress evaporation of the component materials.

Figure 3 is a graph showing the HINAWARI light collector's visible spectrum transmissivity. As can be seen in Figure 3, almost all ultraviolet radiation is excluded, while infrared heat radiation is cut by about forty percent. Light spectrum power analysis was carried out with a Princeton Applied Research Model 1450 OMA with an 0.5 nm wavelength increment.
Gas Exchange System

Figure 6 shows a schematic flow diagram of the system used for measurement of carbon dioxide dissolution into the solution from the gas phase. It uses a hollow fiber module made of micro-porous polypropylene membrane, which was supplied by the Mitsubishi Rayon Co., Ltd.. Figure 7 shows the schematic flow diagram of the system used to recover dissolved oxygen from the solution, which uses the same hollow fiber module. The bore of the hollow fiber was kept at reduced pressure so as to increase partial pressure difference of O₂ over the micro-porous membrane.

Theoretical methods. Carbon dioxide dissolution. The bioreactor contained one liter of pure water, which was circulated through the module at a fixed flow rate (Figure 6). The mixture of air and CO₂ was fed into the bore of the hollow fiber and dissolved into the water flowing outside of the fiber. The concentration of CO₂ in the water, Cₐ, was measured by CO₂ densitometer and recorded. Assuming overall mass transfer coefficient, Kₐ, constant, Cₐ, at any time may be expressed by the following equation:

\[
\frac{C_A - C_{AL}}{C_A} = \exp (-K_a \cdot a \cdot t)
\]

(1)

where

\[
\frac{C_A}{C_A} = H \cdot P_A
\]

(2)

P₀, H, and a are partial pressure of CO₂ in the gas mixture, Henry's constant, and specific membrane area, respectively.

Oxygen recovery. Water dispersed with tiny bubbles of air (instead of oxygen) was fed into the module, where it flowed outside of the hollow fiber. The bubbles attached to the surface of the micro-porous membrane and disappeared gradually; they were sucked through the membrane because the pressure of the bore was kept at about 180 mm Hg. Water did not permeate through the membrane.

Characteristic. Figure 8 shows the effects of concentration of CO₂ on Kₐ at two levels of water flow rate. Figure 9 shows the effects of water flow rate on Kₐ. The values of Kₐ shown in figure were analyzed by Wilson's method to obtain liquid phase mass transfer coefficients, Kₐ, and Kₐ. The overall mass transfer coefficient, Kₐ, in this case may be expressed as follows:

\[
\frac{1}{K_a} = \frac{1}{K_L} + \frac{1}{D} + \frac{1}{H \cdot K_G}
\]

(3)

\[
\frac{1}{K_L} = 1.28 \cdot 10^{-7} \cdot v^{0.6}
\]

(4)

where D/I = 0.248 [m/s]

Where I and D are membrane thickness and diffusivity, respectively, of CO₂ in the gas mixture. V is flow rate of water in terms of m³/s.
EXPERIMENTAL TEST

The bioreactor system consists of the light collection and transmission apparatus, the photoautotrophic bioreactor, and the membrane device for gas exchange (Figure 10). Measurement and control devices were employed for experimentation.

Bioreactor Testing without Gas Exchange System

Apparatus and methods. The light collector, called the "HIMAWARI"/4/, is produced commercially by a Tokyo-based engineering firm, La Foret Engineering and Information Service Co., Ltd. The optical fibers are specially manufactured to be transparent to the visible spectrum (transmissivity of 96% per ten meters), since those used in the communications industry are transparent to the infrared region.

Chlorella was cultivated within the bioreactor in two medium mixtures, one of pH 4.0-4.5 consisting of (mg/L): N, 63.572; P, 227.608; K, 287.308; Na, 4.689; Mg, 394.458; Fe, 0.500; B, 0.500; Mn, 0.500; Co, 0.200; Mo, 9.784, and one consisting of Mayers-4NA5. Inocula were cultivated for two or three weeks in aerated and illuminated (5% CO2 air at 30 L/h and 10,000 lx) glass bottles before being suspended in the bioreactor medium.

Results. Figure 11 shows the effect of temperature of chlorella solution upon the growth rate of chlorella. Although such temperatures are commonly much higher, the optimum growth temperature for chlorella occurred at 24-26°C for this bioreactor at the gas mixture given earlier, perhaps because of the unique nature of visible spectrum light. Figure 12 shows that the rate of chlorella density change increases with density but levels off due to attenuation of the incoming light. Eventually, the rate of density change is adversely affected by increasing the density and an optimum is thus found to be an important operation for minimum energy loss. Figure 13 shows that the rate of chlorella density change changes linearly with respect to the visible light transmission energy. By approximating this linear relationship, higher value data points were plotted.

Fig. 10 Bioreactor system schematic diagram.

Fig. 11 Effect of temperature of chlorella growth rate.

Fig. 12 Relationship between rate of chlorella density change and density of chlorella in basic solution.

Fig. 13 Relationship between rate of chlorella density change and visible ray transmission energy per liter.
Bioreactor Testing with Gas Exchange System

Apparatus and methods. In this testing membrane for gas exchange, the silicone-rubber hollow fiber module was used because this one did not show loadings and a change with passage of operation time. The value of silicone $K_{m} (8.4\times10^{-6} \text{ m/s})$ is higher than polypropylene one. The membrane unit case is made of glass and can be autoclaved. This membrane device was designed by La Fone Engineering, and manufactured by Fuji-Systems Co.. Its specifications are shown in Figure 14.

The operation conditions of gas exchange system are shown in Table 1. Input power levels of the Xenon lamp were 9W, 9W, and 3W in repeated cycles. The incident light intensity in a low earth orbit typically consists of alternating periods of thirty minutes of darkness and sixty minutes of light. Xenon lamps are used as an artificial, controllable source of illumination because of the similarity of their visible spectrum to that of the sun. Xenon lamps coupled with parabolic mirrors to produce a parallel flux can use the same chromatic aberration filtering and fiber-optic ducting as that of sunlight. Because the lamps are separated from the bioreactor by the fiber-optic cables, waste heat is not a concern.

The chlorella solution recycled through the bioreactor and membrane device for gas exchange. The levels of DO and $D_{CO_2}$ were controlled through the pressure and flow rates of the $CO_2$ and $N_2$ supplies. In this test, the level of $O_2$ production could not be analyzed directly. Therefore, the level of $O_2$ production and chlorella production were calculated by the rate of $D_{CO_2}$ change.

Table 1: Comparison between Silicone and Polypropylene.

<table>
<thead>
<tr>
<th>Specification</th>
<th>Operating Condition</th>
<th>Permeability $I$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas exchange membrane</td>
<td>$K_{m} \times 10^{-6} \text{ m/s}$</td>
<td>Effective membrane area $[m^2]$</td>
</tr>
<tr>
<td>Silicone</td>
<td>$8.4$</td>
<td>$0.082$</td>
</tr>
<tr>
<td>Polypropylene</td>
<td>$0.9$</td>
<td>$6.9$</td>
</tr>
</tbody>
</table>

1) membrane thickness 25$\mu$ conversion, $25^\circ C$, $cm^3/m^2.atm.24h$

Results. Figure 15 shows Orbit cycle lights alternation's effect on $D_{CO_2}$ and $DO$ concentration. During the light periods DO concentration increased while that of $D_{CO_2}$ decreased. The opposite occurred during the dark periods. The photosynthetic reaction was demonstrated from Figure 15. During each illumination cycle, the DO and $D_{CO_2}$ concentration pattern was repeated. Figure 16 shows the relationship between visible ray transmission energy and rate of $O_2$ production. The performance of the membrane gas exchanger was constant through the long duration of the experiment.

Fig. 16 Relationship between rate of $O_2$ production and visible ray transmission energy.
CONCLUSIONS

The bioreactor with sunlight supply system and gas exchange system presented here has proved feasible in ground tests and shows much promise for space use as a CEISS device. Our chief conclusions concerning the specification of total system needed for a life support system for a man in a space station are the following (Table 2).

(1) Sunlight supply system: area of solar ray receiving lens: 6.29-9.37 m², electrical power consumption: 37-56 W.
(2) Bioreactor system: volume of chlorella solution: 50 L, radiative surface per liter: 0.58 m²/L, size of bioreactor tank: 0.4 m x 1.0 m.
(3) Gas exchange system: effective membrane area of silicone-rubber hollow module: 1.33 m².

Chlorella photosynthetic efficiency is 7%, which is higher than that one of high plant (0.1-1.0%). Microalgal bioreactor system will be best system for a life support system in a space station.

Table 2 Required performance and simulated specification of total system.

<table>
<thead>
<tr>
<th>Case</th>
<th>I</th>
<th>II</th>
<th>III</th>
</tr>
</thead>
<tbody>
<tr>
<td>Light condition</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Light source</td>
<td>Solar ray</td>
<td>Xenon lamp</td>
<td>Solar ray</td>
</tr>
<tr>
<td>Total mass of O₂ produced per 1 day in tank [g/day]</td>
<td>686*1.1=754.6</td>
<td>686*1.1=754.6</td>
<td>686*1.1=754.6</td>
</tr>
<tr>
<td>Average mass of O₂ produced per hour during light time [g/h]</td>
<td>31.4</td>
<td>31.4</td>
<td>47.2</td>
</tr>
<tr>
<td>Chlorella biomass produced per hour [g/h]</td>
<td>29.5</td>
<td>29.5</td>
<td>44.2</td>
</tr>
<tr>
<td>Rate of chlorella density change [g/L/h]</td>
<td>0.59</td>
<td>0.59</td>
<td>0.88</td>
</tr>
<tr>
<td>Volume of chlorella solution [L]</td>
<td>50</td>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td>Radiative surface per liter [m²/L]</td>
<td>0.58</td>
<td>0.58</td>
<td>0.58</td>
</tr>
<tr>
<td>Visible ray transmission energy per liter needed in tank [kWh]</td>
<td>49a)</td>
<td>49a)</td>
<td>72b)</td>
</tr>
<tr>
<td>Temperature range of solution [°C]</td>
<td>25±1</td>
<td>25±1</td>
<td>25±1</td>
</tr>
<tr>
<td>Density range of chlorella in solution [g/L]</td>
<td>about 10a)</td>
<td>about 15b)</td>
<td>about 15b)</td>
</tr>
<tr>
<td>Chlorella photosynthetic efficiency [%]</td>
<td>7</td>
<td>7</td>
<td>7</td>
</tr>
<tr>
<td>HIMWARI transmission efficiency [%]</td>
<td>60</td>
<td>-</td>
<td>60</td>
</tr>
<tr>
<td>Area of solar ray receiving lens [m²]</td>
<td>6.29</td>
<td>-</td>
<td>9.37</td>
</tr>
<tr>
<td>Visible ray transmission energy [kW]</td>
<td>2450</td>
<td>2450</td>
<td>3650</td>
</tr>
<tr>
<td>Electrical power consumption [kW]</td>
<td>37</td>
<td>24500</td>
<td>56</td>
</tr>
<tr>
<td>Gas exchange module</td>
<td>Silicone-rubber hollow fiber</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Effective membrane area [m²]</td>
<td>Max. 0.082*50[L]/3.1[L] = 1.33</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Size of module [mm]</td>
<td>145*450</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

a) experimental data  b) predicated data

REFERENCES

2. K. Mori, Photoautotrophic bioreactor using visible solar rays condensed by Fresnel lenses and transmitted through optical fibers, Biotechnology and Bioengineering, No.15, p.331 (1986)

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