Assessment of AMBR™ as a model for high-throughput cell culture process development strategy

Sarath Moses1*, Matthew Manahan1, Alexandre Ambrogelly2, Wai Lam W. Ling1*

1Upstream Process Development & Operations, Bioprocess Development, Merck Research Laboratories, Union, USA
2Analytical Development, Bioprocess Development, Merck Research Laboratories, Union, USA
Email: *sarath.moses@merck.com, wai_lam.ling@merck.com
Received 20 August 2012; revised 27 September 2012; accepted 11 October 2012

ABSTRACT

The development and delivery of high quality therapeutic products necessitates the need for high-throughput (HTP) process development tools. Traditionally, these works require a combination of shake flask and small-scale stirred tank bioreactor (STR) which are labor and resource intensive and time-consuming. Here we demonstrate a strategy for rapid and robust cell culture process development by evaluating and implementing the use of a new HTP disposable micro bioreactor (MBR) called AMBR™ system (Advanced Microscale Bioreactor) that has the capabilities for automated sampling, feed addition, pH, dissolved oxygen (DO), gassing and agitation controls. In these studies the performance of two monoclonal antibody (MAb) producing cell lines (MAb1 and MAb2) was evaluated both in the AMBR system and 3-L STR. We demonstrated that cell culture performance (growth and viability, production titer and product quality) were similar in both vessel systems. Furthermore, process control and feed optimization were demonstrated in an additional cell line (MAb3) in the disposable MBR and its performance confirmed at STR scale. The results indicate that the AMBR system can be used to streamline the process development effort and facilitate a rapid and robust cell culture process development effort for MAb programs in a HTP manner.

Keywords: Microbioreactors, High-Throughput, DoE, Process Ranging, Process Development, Cell Culture

1. INTRODUCTION

The continuing trend in biopharmaceutical industry in delivering quality therapeutic products into the market with shrinking timelines and limited resources has facilitated the development and implementation of HTP systems to enhance data outputs and program decision in the cell culture process development area. In addition, recent ICH (International Conference on Harmonisation of Technical Requirements for Registration of Pharmaceuticals for Human Use) endorsed guidelines for ICH Q8/Q9/Q10/Q11 implementation, [1] focuses on ensuring the quality of the therapeutic products to be safe and efficacious, and provides guidance on risk assessment and management on product quality control through identification of critical quality attributes (CQAs) and critical process parameters (CPPs). Our ongoing efforts in applying Quality by Design (QbD) approaches in high throughput cell culture systems are one of the strategies in assessing the product quality changes during development while balancing the resource and timeline constraints.

The ongoing efforts in the biopharmaceutical industry to balance the increased workload required to address process and product quality control coupled with the reduced resources availability have heightened the implementation of HTP systems to support development work. One of the HTP strategies utilized in upstream process development is the use of disposable micro-bioreactors for cell culture process development. These disposable micro-bioreactors are expected to perform in a high-throughput manner under controlled bioreactor operation conditions similar to those at bench and large-scale, but without extensive bioreactor set-up and clean-up procedures. These systems may replace the use of shake flasks for early stage of cell culture process development and optimization studies [2-5].

Recent literature have demonstrated availability of various designs of micro-scale reactors from a simpler standard plate with integrated sensors called Microtiter plate based bioreactors (MTB) [6-12] to more advanced cell culture systems such as the Stirred Mini Bioreactors (MBR) with control capabilities close to bench-top bioreactors for high throughputs cell culture screening and optimization studies [13-17]. MBRs possess the advantage of being able to closely mirror the bench-scale bio-
reactors in their monitoring and control capabilities and hence are considered as a preferred alternative vessel over shake flasks or MTBs for early stage process development. MBRs are generally equipped with optical probes to monitor pH and DO online and it has been extensively characterized for its oxygen transfer rate, mixing capabilities and gas sparging efficiency [13-17]. The above mentioned characteristics of MBRs typically make them an ideal set up for early high-throughput clonal screening efforts as part of process development.

In order to assess our HTP cell culture process development needs the ideal system must address the following requirements: 1) be able to individually control pH and DO; 2) have the capability for automated medium, feed and base additions; 3) have reduced operational requirement compared to standard STR, 4) utilize disposable vessels, thus minimizing the cleaning and sterilizing procedure for a quick turn-around time; and 5) relatively high number of available vessels to be run and managed simultaneously which provide needed statistical significance for process parameters under. In this manuscript, we present the results of the study evaluating and implementing the new disposable MBR called AMBR system manufactured by The Automated Partnership (TAP Biosystems, UK) in a head-to-head comparison with 3-L bench-top STR. For this study, we compared two established MAb production processes (MAb1 & MAb2). The study assessed the comparability of process parameters such as pH, temperature, dissolved oxygen (DO) and osmolality between the AMBR and 3-L STR and their overall impact on cell culture performance, metabolic profiles and product quality. Furthermore, the AMBR system was utilized in a central composite DoE (Design of Experiment) based pH & DO optimizations study for a MAb (MAb3) cell culture production process, and the cell culture performance results from the optimized process was confirmed and reproduced in 3-L STR.

2. MATERIALS AND METHODS

2.1. Media, Cell Lines and Culture Propagation

Animal component-free growth medium was purchased from SAFC (St. Louis, MO). All feed media were developed in-house containing no animal-derived raw material and manufactured by SAFC and Hyclone (Logan, UT). Suspension adapted Chinese Hamster Ovary (CHO) cell lines expressing 3 different monoclonal antibodies MAb1, MAb2 and MAb3 were used. These MAb production cell lines were serially sub-cultured in growth medium. For MAb1 and MAb2 cell lines the cells were maintained in shake flasks in a 37°C, humidified 7.5% CO2 incubator with agitation at 100 rpm. The typical cell inoculation density was 3.0 - 5.0 × 10^5 cells/mL and the culture was routinely passaged three times per week. For MAb3 cell line the cultures were maintained and propagated as described in [9].

2.2. Bioreactor Systems

AMBR-24 is an automated MBR system with 24 disposable microbioreactor vessels. The AMBR instrument is housed in a bio-safety cabinet to facilitate sterile environment for sampling and feeding. The system includes two culture stations CS-1 and CS-2 containing 12 microreactors each. Each microbioreactor vessel has a miniature internal impeller, as well as individual closed loop control of dissolved oxygen, pH and gas supply. Each culture station has independent stirring speed and temperature control, allowing up to two different temperature and stir speeds simultaneously. AMBR system can be programmed for automated culture preparation and inoculation, feed and base addition, and culture sampling scheduling necessary for cell count, metabolite and latter analysis. Monitoring of pH and DO is via optical sensors embedded in disposable microbioreactor vessels. The measurements are performed at a frequency of every 90 seconds by the sensors.

The 3-L STR (B. Braun Biotech, Bethlehem, PA, Sartorius) were operated with a 2-L working volume. Temperature was controlled using an electric heating blanket at 37°C. The pH was maintained using a cascade of CO2 and sodium hydroxide additions. Dissolved oxygen (DO), measured as percent air saturation at 1 atm pressure, was maintained at set point by intermittent oxygen sparging, constant air flow in the headspace, and constant agitation. The cultures were mixed using dual pitched-blade impellers.

2.3. MAb1 and MAb2 Cell Culture Process

Cell inoculum scale-up was prepared in shake flasks at seeding density of 3.0 - 5.0 × 10^3 cells/mL for both the MAb cultures and ready for inoculation into production vessels 2 - 3 days after initial seed. Fed batch studies in 3-L STR (2L working volume) and in AMBR (12 mL working volume) were seeded with cells at ~3.0 × 10^5 cells/mL obtained with a split ratio of 1:4. Samples were taken every 2 - 3 days to monitor for cell growth, metabolites, and MAb production. Glucose and glutamine were supplemented as needed to avoid depletion.

For initial comparison for MAb1 and MAb2, the evaluation in AMBR system was performed in replicates of 12 for each MAb, with 6 disposable microbioreactor vessels in each culture station. The positions of the MAb cell lines are depicted in (Figure 1).

Cell culture temperature was maintained at 37°C for the first three days and then shifted to 34°C for the remaining of the run. Unless indicated, media pH was set.
Culture Station 1

Culture Station 2

MAb1

MAb2

Figure 1. Schematic representation of AMBR culture station and the experimental design setup of MAb1 and MAb2 cell line process. Each process was run in replicates of 6 in each of the culture stations.

at 6.8 with pH deadband at 0.05, DO at 60% for both 3-L and AMBR bioreactor systems, and agitation rate was set at 200 rpm for 3-L STR and 900 rpm for AMBR (per manufacturer's recommendation) to maintain a tip speed of 0.50 m/s. pH of the culture was controlled by addition of 1M NaOH or sparging CO2 gas. Antifoam was added as needed during the run. For the 3-L STR, DO level was maintained at the set point with air overlay or oxygen delivered from the sparger.

2.4. MAb3 Cell Culture Process

Cell inoculum scale-up was prepared in shake flasks at seeding density of 2.0 - 4.0 × 10^5 cells/mL and ready for inoculation into production vessels 3 - 4 days after initial seed. Fed batch studies in 3-L STR and in AMBR were seeded with cells at ~2.0 × 10^5 cells/mL. Culture temperature was maintained at 37°C throughout the process. Media pH was set at 6.9 with pH deadband at 0.05, DO at 30% for both bioreactor systems. Agitation rate was set at 200 rpm for 3-L STR and 900 rpm for AMBR. Samples were taken every 2 - 3 days to monitor for cell growth, metabolites, and MAb production. Glucose was supplemented as needed to avoid depletion.

2.5. Cell Culture Analyses

Cumulative viable cell mass, CVC, was calculated by the following equation:

\[
CVC(t) = \int_0^t X_v(x) \, dx \approx \sum_{i=1}^{N} (t_i - t_{i-1}) \left( X_{v,i} + X_{v,i-1} \right) / 2
\]

Where, \(X_v(x)\) is the viable cell density at time \(x\), \(X_{v,i}\) is the viable cell density at time \(t_i\), \(X_{v,i-1}\) is the viable cell density at time \(t_{i-1}\). CVC \((t)\) is the cumulative viable cell density at time \(t\). The specific production rates of product were calculated as the linear regression line of the product concentration versus CVC. Statistical analyses were performed using SAS-JMP software 8.0 (SAS North Carolina, USA).

Specific glucose consumption, lactate production and lactate consumption rates were estimated by plotting cumulative glucose or lactate versus the CVC and fitting the plots with a regression coefficient of close to one. The slope of this line was used as an average specific rate.

2.6. Analytical Methods for Cell Culture and Product Quality Analysis

Viable cell density (VCD) and total cell density in shake flasks and bioreactors were measured by CEDEX (Roche USA, Indianapolis). Glucose, lactate, glutamine, and glutamate were determined by YSI 2000 (YSI, Yellow Spring, OH). Osmolality was detected by Advanced Micro-Osmometer (Advanced Instruments, Norwood, MA). pH, pCO2, pO2 were measured by ABL5 (Radiometer America Inc, Westlake, OH).

MAb production in bioreactors was determined by Reversed Phase HPLC (RP-HPLC, Poros R2/10, Applied Biosystems). CVC and Qp were calculated using cell density values determined by CEDEX and MAb titers by RP-HPLC [18]. Product quality of Protein-A purified intermediates was analyzed by high performance size exclusion chromatography (HP-SEC, GE Healthcare Superdex 200 10/300 GL column), ion-exchange chromatography (IEX, Dionex Propac WCX-10 4 x 250 mm column) and reversed-phase HPLC (RP-HPLC, Poros R2/10, Applied Biosystems).

3. RESULTS

3.1. Assessment of Performance between AMBR and STR

3.1.1. Process Parameters

To ensure that this MBR system can mimic DO, pH,
temperature controls observed for the 3-L STR, online profiles were collected and showed comparability between the 2 bioreactor systems (Figure 2). Dissolved oxygen level of MAb1 and MAb2 process was measured.
Figure 2. Online process parameter profile in AMBR and STR for both MAb1 (a), (c), (e) and MAb2 (b), (d), (f) cell lines. DO (a), (b), pH (c), (d), and temperature (e), (f). (▲) AMBR CS-1, (●) AMBR CS-2, and (○) STR. The mean of all online values (n = 6) of the culture vessels from the corresponding culture stations are plotted alongside the value of 3-L STR (n = 1).
online for both the AMBR and 3-L STR over the period of the process run. In MAb1 cell line the set point DO of 60% was maintained by AMBR vessels as detected by the vessel biosensors in both culture stations, however in 3-L STR the trend was maintained between 50% and 80% as detected by the DO probe. In MAb2 cell line the DO was maintained at the 60% set point in both AMBR culture stations and 3-L STR vessel. pH values of MAb1 and MAb2 process was measured over the duration of the cell culture runs online in AMBR and 3-L STR (Figures 2(c) and (d)). The pH set point was at pH 6.8 and dead band at 0.05. Online values showed an acceptable variation range of ±0.05 from the set point between AMBR and 3-L STR and also between individual reactor vessels in each of the culture stations [17]. One of the disadvantages of earlier version of pH optical sensors utilized in other MBR was the loss of sensitivity to pH change with time, leading to the inability of these vessels to control pH later in culture [3,15]. In these demonstrated runs using the AMBR vessels embedded with optical pH sensors, no significant variations in pH readings were identified in all 24 culture vessels of the AMBR system over the 19 day of culture duration.

Each of the AMBR culture station blocks CS-1 and CS-2 can be maintained independently at different temperatures. Temperature control in AMBR is influenced by the environmental temperature [17] and care should be taken that it is maintained in an ambient condition between 21°C - 25°C as per manufacturer’s instruction. In this study AMBR measurement of culture temperature online demonstrated temperature control to be consistent with that of the 3-L STR as shown for both MAb1 and MAb2 processes (Figures 2(e) and (f)). Temperature control between the culture stations 1 and 2, as well as amongst each of the culture vessels within the stations was consistently maintained. Temperature shift was performed from 37°C to 34°C on day 3 in both the systems and maintained till the end of the run. One of the primary advantages of the AMBR over other known MBR systems is that the AMBR does not require agitation to pause for sampling or feed addition hence reducing interruption in temperature maintenance [14]. The mixing time of the feeds were also not impacted since the agitation is not affected. This characteristic feature of temperature stability in AMBR helps maintain the overall cell culture process reproducibility which can be replicated in large scale cell culture where culture temperature is not typically impacted by feed additions.

The offline trend in osmolality between the MAb1 and MAb2 process in both the AMBR and STR were shown to be similar. For MAb1 process the osmolality gradually increased from approximately 300 to 350 from day 0 to day 7 and subsequently return to 300 by the end of the culture run (Figure 3). For MAb2 process the offline osmolality peaked on day 6 to approximately 375 and decreased to 350 by the end of the culture run. The comparable process parameter results detected in 2 individual cultures with independent processes from the AMBR and the 3-L STR demonstrate the AMBR has the potential to replicate the performance of bench-scale bioreactors.

3.1.2. Cell Culture Parameters
VCD and viability of both MAb1 and MAb2 cell lines were analyzed offline using CEDEX at regular intervals during the process and showed consistency between the AMBR stations and 3-L STR throughout the culture runs (Figure 4(a)). A peak cell density of approximately 4.0 - 5.0 × 10^4 cells/mL and 6.0 - 7.0 × 10^4 cells/mL was achieved in both AMBR and 3-L STR on day 10 for MAb1 and MAb2 cell lines respectively. Viability was maintained at ~95% until day 12 for both MAb1 and MAb2 in AMBR and 3-L STR after which it trended down gradually to 80% for MAb1 and 60% for MAb2 cell lines at the time of harvest (Figure 4(b)). The plot also shows overall consistency in VCD and viability between culture stations in AMBR in each of the processes.

Protein production in both MAb1 and MAb2 cell culture process followed a consistent trend between AMBR and 3-L STRs. The final titer at the end of the production batches were approximately 1.2 ± 0.2 and 1.0 ± 0.1 arbitrary units for MAb1 and MAb2 respectively in both AMBR and 3-L STRs (Figure 4(c)). More importantly, protein quality attributes of Protein-A purified MAb from the AMBR and STR systems determined by RP-HPLC, HP-SEC (Figure 4(d)) and IEX (Figure 4(e)) indicated comparable product quality profiles.

The metabolic profile for both MAb1 and MAb2 cell lines were measured and represented as rates (Table 1).

![Figure 3. Offline measurement of Osmolality for AMBR and STR in both MAb1 and MAb2 processes. (a) MAb1:AMBR CS-1; (●) MAb1:AMBR CS-2; (▲) MAb1:STR; (○) MAb2:AMBR CS-1; (●) MAb2:AMBR CS-2; and (▲) MAb2:STR.](http://www.sciencedirect.com/science/article/pii/S095340951200177X)
S. Moses et al. / Advances in Bioscience and Biotechnology 3 (2012) 918-927

Figure 4. Offline monitoring of (a) viable cell density, (b) cell viability, (c) MAb titer. (▲) MAb1: AMBR CS-1; (●) MAb1: AMBR CS-2; (●) MAb1: STR; (▲) MAb2: AMBR CS-1; (●) MAb2: AMBR CS-2; and (○) MAb2: STR. Product quality assessment for MAb1 and MAb2 cell culture processes in AMBR and STR by (d) RP-HPLC and HP-SEC (e) IEX. The mean of all online values (n = 6) of the culture vessels from the corresponding culture stations are plotted alongside the value of 3-L STR (n = 1).

Table 1. Metabolic profiles for MAb1 and MAb2 processes between AMBR and STR.

<table>
<thead>
<tr>
<th>Bioreactor</th>
<th>Glucose Consumption Rate*</th>
<th>Lactate Production Rate†</th>
<th>Lactate Consumption Rate*2</th>
</tr>
</thead>
<tbody>
<tr>
<td>MAb1 Process</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AMBR CS-1</td>
<td>–428.8 ± 6.0</td>
<td>218.2 ± 5.7</td>
<td>–21.8 ± 1.5</td>
</tr>
<tr>
<td>AMBR CS-2</td>
<td>–404.1 ± 4.8</td>
<td>221.1 ± 6.0</td>
<td>–21.1 ± 1.1</td>
</tr>
<tr>
<td>3-L STR</td>
<td>–447.1</td>
<td>162.4</td>
<td>–18</td>
</tr>
<tr>
<td>MAb2 Process</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AMBR CS-1</td>
<td>–356.4 ± 4.4</td>
<td>242.3 ± 14.7</td>
<td>–14.6 ± 2.4</td>
</tr>
<tr>
<td>AMBR CS-2</td>
<td>–347.9 ± 12.3</td>
<td>245.6 ± 36.2</td>
<td>–14.2 ± 0.7</td>
</tr>
<tr>
<td>3-L STR</td>
<td>–478.0</td>
<td>220.4</td>
<td>–11.6</td>
</tr>
</tbody>
</table>

*All rates are represented as pg/cell-day; †n = 6; ‡n = 1; †Lactate production during early production phase; ‡Lactate consumption during later production phase.

The initial consumption rates of glucose for MAb1 process were similar across the different culture stations of the AMBR and 3-L STR. However the glucose utilization rate was higher for the MAb2 process in 3-L STR at ~478 pg/cell-day compared to those from the AMBR culture stations with an average rate of ~350 pg/cell-day.

The lactate production rates for MAb1 process in 3-L STRs (~162 pg/cell-day) showed a decrease of ~26% compared to the two culture stations in AMBR (average...
~219 pg/cell-day). Conversely the lactate production rates for MAb2 process were similar between the AMBR and 3-L STRs at ~244 pg/cell-day and ~220 pg/cell-day, respectively. The lactate consumption rates during the later stage of the cell culture runs followed similar trending of lower consumption rates resulted from lower production rates for both MAb1 and MAb2 processes.

3.2. Optimizing the Effect of pH and DO of an MAb Process Using AMBR

The comparable online profiles of pH and DO demonstrated for MAb1 and MAb2 between the STR and the AMBR provided confidence that initial pH and DO process optimization can be performed in a high throughput manner using the AMBR. For MAb3, an initial pH and DO screening was conducted with the AMBR to enhance MAb3 productivity (Figure 5). Multivariate analysis (Figure 5(a)) for pH, DO and titer demonstrated that pH set point had a significant effect on MAb titer (p < 0.05) while DO did not (p > 0.05). Regression analyses of specific productivity (Qp) as a function of pH setpoints demonstrated positive linear relationship (Figure 5(b)). The increase in titer productivity could also be attributed to the increase in the CVC at higher pH ranges (Figure 5(c)). Overall pH 7.0, the highest pH setpoint tested and DO at 30% were determined to be the optimal process condition for further confirmation in 3-L STR.

To assess the scalability of the AMBR optimized cell culture process, 4 replicate runs were conducted in 3-L STR with chosen setpoints of pH at 7.0 and DO at 30%. As shown in (Figures 5(d) and (e)), the cell culture performance in 3-L STR as represented by CVC and titer productivity of MAb3 cell line process was found to be comparable and more importantly scalable with the data obtained from AMBR.
4. DISCUSSION & CONCLUSION

Cell culture process development timeline is driven by historical experience and existence of platform. However, new MAb molecules entering the development pipeline are required to not only meet specific product quality profiles, but also develop in shorter timelines. As a result, high throughput screens to develop production processes become critical path to meet these demands. The emergence of single use disposable bench-scale and large-scale bioreactors has facilitated rapid development and manufacturing of drug supplies. We demonstrate a similar approach is applicable for high throughput process development. While disposable bench-scale bioreactors can reduce set-up and break-down time, no time or resource saving was apparent in cell inoculum and medium preparation or batch operation. We have selected the use of AMBR system fitted with disposable microbioreactors for process development, as they have low inoculum and medium requirement and thus minimize operational workload and improve throughput. The system also has a well integrated technology for automated inoculation, feeding and sampling. The disposable nature of the AMBR vessels reduces vessel set-up and clean-up effort.

Based on our studies, the AMBR system is shown to be a versatile platform for HTP process development and optimization for mammalian cell cultures. We have demonstrated that the AMBR is an excellent and efficient tool to support a number of HTP DoE-based process development investigations. These include identification of feed medium concentrations, pH and/or DO optimization ranges for a mammalian cell line, comparison of culture performance of multiple cell lines with similar processes, or conducting process ranging studies. The utilization of HTP DoE-based investigation is most advantageous during initial stage of process development as this system enhances data outputs with reduced timeline, and thus allows program decision for cell culture area to proceed earlier. Nevertheless, early process ranging studies conducted using this HTP DoE-based approach to narrow the process test ranges would significantly reduce the more labor intensive, subsequent work in small scale STR.

While our results are supportive of the AMBR in meeting most of the STR capabilities, there are several points which would allow AMBR or other MBR systems to achieve more desirable operational controls and the potential to be a scale-down model for large-scale vessels. First, continuous feed strategy cannot be achieved. An attempt to design a continuous feed strategy with the MBR will require multiple and frequent small feed aliquot additions. Since this MBR system is not a closed system, introduction of frequent feed additions would change the gassing regime and may affect the overall pH and DO of some cell cultures. Second, continuous sparging in the MBR necessitates frequent antifoam additions that may not be required for some cultures in standard STR. Third, culture temperature control in the MBRs like AMBR can be affected by environmental temperature. Fourth, the current state of this MBR vessel and impeller configurations are different from those of the standard STR, and thus processes developed and KLa calculated using this MBR may not be scalable to larger vessels for some cultures. With further development and design modification these limitations could be addressed in the future.

REFERENCES


