Mixing Performance Analysis of Orbitally Shaken Bioreactors

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Featured Application: Orbitally shaken bioreactors (OSBs) are the most frequently used type of bioreactor in the synthesis of polyhydroxyalkanoates from bacteria or algae or the extraction of biomolecules from biomass for bioprocess development. Their success can be attributed to their simple and functional design, which make shaking systems suitable for a large number of cost-efficient parallel experiments. Recently reported findings for power-input, in-phase and out-of-phase operation and mixing in OSBs are summarized in this article. Novel mixing-time monitoring techniques for the control of mixing conditions in shake flasks, Erlenmeyer flasks and microwell plates are described. The methods for characterizing mixing conditions and the novel mixing time online measurement techniques that are summarized in this article can be utilized to tap the full potential of OSB systems. A new set of correlations for the Newton (power) number and mixing time when liquid motion was induced in the OSB that can be considered as being fully validated on scale-up.

Abstract: This study investigated the efficacy of a novel correlation of power input, energy dissipation rate and mixing time as a potential route to identify the orbitally shaken bioreactor (OSB) system. The Buckingham’s π-theorem was used to designate and transform dimensionless Newton numbers with five relevant power input variables. These variables were empirically varied to evaluate the correlation among the dimensionless numbers. The Newton number decreases with the increased shaking frequency and filling volume. Previous work has focused on optimizing the mixing process by evaluating different shaking and agitation mixing methods. We establish a new mixing process and assessable measurement of the mixing time in the OSB. An innovative explanation of mixing time for the thermal method is proposed. The optimal mixing time is independent of the temperature of filled liquid. The dimensionless mixing number remained constant in the turbulent regime and increasing with the increased liquid viscosity and filling volume. Our findings revealed that the observed correlation is a practical tool to figure the power consumption and mixing efficiency as cell cultivation in all OSB scales and is fully validated when scaling-up system.

Keywords: orbital shaken; bioreactor; mixing time; power consumption

1. Introduction

Mammalian cell cultures play a crucial role in the biopharmacy industry and are an efficient way to obtain pharmaceutical protein, which is in increasing demand. Bioreactors are an important piece of equipment for mammalian cell cultivation, which indicates that the development of the bioengineering
industry is largely determined by bioreactors capability. There are several types of bioreactors with different agitating principles. Stirred-tank bioreactors (STRs) are the most traditional bioreactor design, which has been fully developed and its largest scale can be up to 20,000 L. Air lift bioreactors employ moving gas bubbles with a high speed to agitate the fluid in the bioreactor, which can produce a high shear stress resulting in mammalian cell death. Wave bioreactors can mix the fluid efficiently, but this type of bioreactor is difficult to scale up due to the cumbersome mechanical system and its largest scale is only about 500 L. Recently, orbitally shaken bioreactors (OSRs) have become increasingly popular in processing cell cultures because of its simple agitating principle, low-cost, ease of operation and suitability for single use experiments. Nowadays, the CHO cells cultivation with 2000 L in OSRs was reported and this type of bioreactor is considered as the most promising candidate to take the place of STRs. Prior to their general adoption by industry, large-scale industrial mixing tanks were used in the early stages of bioprocess development, such as microbial fermentation, biotransformation and recycling technologies. A microscale method depends on the utilization of shaken microtiter plates. On a shaken microtiter plate, full mixing is accomplished by shaking, rather than stirring. Shake flask bioreactors are cost-effective and simple to operate. They are available in 25-mL to six-liter volume sizes, different geometries and materials of construction [1], but limited to batch system, mass transfer, scale-up and real-time monitoring [2]. Platas et al. [3] identified the specific growth rate to shake flask process parameters of specific power consumption, mixing time, maximum fluid velocity and Reynolds number. High-throughput shaking bioreactors are more flexible than conventional stirred-tank bioreactors, which are less automated and restricted to off-line sampling due to their small volume capacity [4]. Table 1 summarizes the relation between the experimental throughput and the level of monitoring and control required for an effective cell cultivation process. Mixing and oxygen-mass transfer are both enhanced by shaking. The rate of both processes is influenced by the geometry of the flask, the fill volume and the intensity of shaking (frequency and amplitude of shaking) [5]. Compared to conventional stirred-tank bioreactors [4], the advantages of shaking bioreactors are ease of operation, lower material requirement and capital investment, higher throughput and lower labor requirement. However, the disadvantages are lower oxygen-mass transfer efficiency, less accessible to monitoring and control, scale-up criteria not well established and lower information content per experiment.

### Table 1. Summarized conventional and high-throughput equipment [4–7].

| Type                      | Volume       | Typical Experimental throughput | Level of Monitoring | Cost of Operation                        |
|---------------------------|--------------|---------------------------------|---------------------|------------------------------------------|
| Microtiter plates         | 100 µL–3 mL  | Very high: thousands per technician | Low: T, biomass, product | Medium: capital, increased use of disposables |
| Small-scale stirred vessels | 10 mL–100 mL | Low/medium: 20 per technician | Low: T, biomass, product | Low |
| Miniaturized stirred bioreactor | 25 mL–6 L. | Medium: 50 max. per technician | High: pH, oxygen, T, biomass and product | Medium: capital, labor |
| Shaken vessels            | 2 L–50 L.   | Low: 1–5 technicians            | High: pH, oxygen, T, biomass and product | High: capital, raw materials, labor |
| Conventional stirred-tank bioreactor | 1 L–100 L. | Low: 1–5 technicians            | High: pH, oxygen, T, biomass and product | High: capital, raw materials, labor |

Shaken bioreactors were comprehensively utilized in the early stages of bioprocess development and microscale plates (24-, 48-, 96- or 384-orifice discs) facilitated a significant number of experiments. The shaker (25 mL–6 L) was the most widely used orbitally shaken bioreactor (OSB) type, providing a cost-effective system that is effective for a broad range of applications [8–14]. Orbital-shake bioreactors are universally utilized for small-scale screening and process adjustment. Complicated mechanical and electronic components of the shaken table drive, the power supply and the control unit are integrated into the shaken plate and are therefore independent of the reactor. This makes the reactor design simple and cost-effective, not only for many small-scale parallel experiments, but also for general one-time applications. Klöckner et al. [15] proposed a simplified design, reducing complexity and improving
the accessibility of serviceable parts such as rotating impellers and baffles. Klöckner and Büchs [6] demonstrate adequate performance with regards to measured oxygen transfer capacity values for larger than 500 rpm high shaking frequencies and three and six-millimeter-low shaking diameters, which were conducted in different microtiter plate materials in combination with polystyrene and deionized water.

The design of the bioreactor is divided into two aspects, namely, power consumption and mixing efficiency. Power consumption is the most essential significant parameter describing flow regime and mass transfer characteristics in designing and operating bioreactors. The local power consumption of the reactor refers to the external input energy required to keep the fluid motion, and the magnitude of the local power consumption represents the power output of the power system of the reactor in each case. In addition, local power consumption can also reflect the intensity of fluid motion inside the reactor to a certain extent. In general, the more intense the fluid movement, the stronger the material mixing uniformity, but the greater the fluid shear force, increasing the risk of cell death. Therefore, the analysis of the local power consumption of bioreactors can often make a comprehensive analysis of cell growth metabolism problems such as substance mixing and cell damage. The power is transmitted during the shaking process, and different measurement systems are described to determine the average unit volume power ($P/V_L$) of the cylindrical bioreactor. The specific $P/V_L$ is one of the basic parameters for microbial process scale-up and scale-down. Parameters that directly affect biological culture, such as oxygen supply, mixing and dispersion of hydraulic stress or organic fluid phases are reliant on specific power inputs [1]. The consequences of changing flask size, shaking frequency and filling volume in relation to observed viscosity change over time during the fermentation process is well documented [15,16]. During the fermentation process viscosity of the ferment increases, necessitating an increase in power input to maintain consistent mixing [17]. The $P/V_L$ values in the cylindrical shaken disposable (single-use) reactor ranged from 0 to 2 kW/m$^3$ which depend on liquid viscosity, the filling quantity and shaking frequency. Klöckner et al. [18] introduced current power input correlations as a possible way to describe for the hydrodynamics of the news systems described above. A set of related power-consumption variables is demonstrated with Buckingham’s $\pi$-theorem and converts them to dimensionless Newton numbers. The related numbers are experimentally altered to evaluate the relationship between Reynolds and Froude numbers. A simple correlation is developed for $P/V_L$ with seven variables to verify with the 200 L and 2000 L orbitally shaken bioreactors application. During cell culture fermentation at all scales, the power input and hydrodynamics for cylindrical orbitally shaken bioreactor systems can be predicted using this effective, established correlation tool.

Mixing efficiency is a key engineering aspect of the bioreactor because it ensures the homogeneity of the culture ferment [19]. An understanding of the characteristics of mixing are therefore necessary to determine any regions of heterogeneity within the bioreactor and gauge resulting concentration gradients that cells may encounter. For the direct method of mixing efficiency determination used the measurement of the mixing time in the bioreactor, which is designated as the time desired to accomplish a certain degree of homogeneity in the liquid [20]. Mixing time measurement methodologies can be distinguished into local and global technologies by the measured volume. The local methods relied on physical measurements (pH, temperature or conductivity) with intrusive probes at specific point. The global approach is not intrusive and depends either on chemically based reactions and/or optical measurement of different mixing regions in the reactor [21–23]. However, global approaches are usable only in transparent vessels and are less accurate than local methods.

Exact mixing time measurement is essential to minimize the concentration gradient that will be harmful to the cell culture in a shaken reactor [24]. Several different methods have been developed for the measurement of mixing time: conductivity or pH method, thermal method, color addition or decolorization methods [25,26]. Tan et al. [26] utilized the colorimetric method to measure the mixing time. The reported mixing time was the same for both 25- and 50-mm shaken-flask diameter. The dimensionless mixing number remains constant and is independent of the given geometry of the flasks under high Reynolds numbers region. Tan et al. [26] utilized a single indicator system to gauge
the mixing time in the Erlenmeyer shaken flasks. Tissot et al. [27] used a two pH indicators technique to measure the mixing time in the shaken cylindrical vessels, concluding that the orbital diameter had no symbolic effect on the mixing time in the shaken flask. The fastest diffused mixing area was determined to be in the lower region of the flask. At a low shaking frequency, two different zones could be divided inside the reactor. The first zone is close to the free surface where mixing was increased by convection, and the second zone is in the lower region of the cylinder where mixing was driven by diffusion. Rodriguez et al. [28,29] compared the orbitally shaken bioreactor and the Erlenmeyer flask with a stirred tank, measuring mixing time and classifying two mixing mechanisms; in-phase and out-of-phase. Mixing time was accurately determined using a dual indicator system (DISMT) to optimize and minimize concentration gradients for harmful cell cultures prevention. For more efficient mixing, the feeding pipe was inserted near the reactor wall and the outside of the vertical structure close to the center of the reactor. Weheliye et al. [30] indicated the correspondence of the flow transition phenomena at 10-cm inner diameters and 13-cm fluid heights different cylindrical bioreactors. The results also show the feed-pipe position affected the mixing time in the cylindrical reactor. For both systems, it is suggested to insert the pipe near walls with higher shear stress close to the boundary layer of the wall. The highest/lowest sides of the torus are regularly coordinated with the centrifugal acceleration of the shaker table that shifts its direction as phase angle of table increases. The two vortices expand over the whole cylinder height and finally control the flow inside the bioreactor at 110 rpm shaking frequency.

Hoogendoorn and Hartog [25] studied the batch mixing of viscous liquids with rotary mixers in 0.24 and 1.8 m diameter vessels. Power input and mixing time were the measured parameters. Mixing time measurements were made using discoloration and thermal-response techniques, both of them generating reproducible, quantitative data. For some agitators in the real viscous flow region, the mixing time is inversely proportional to the agitator speed. The mixers could be compared in terms of power number and mixing number based on liquid viscosity and density, power consumption, mixing time and vessel diameter. Havas et al. [31] used the conductivity method, thermal method and color addition method to measure mixing time. Experiments were executed in a 396-mm-diameter tank with basket-curve bottom and a 529-mm-diameter tank with plain bottom. They found the most suitable method according to thermal pulse disturbance. The mixing number is the product of mixing time and the rotation speed of the agitator. Neither the temperature and the place of thermal pulse disturbance, nor the location of the measuring point has a significant effect on the mixing time which the dimensionless mixing number is independent of high Reynolds number (Re > 10^6) within the turbulence range [31,32]. The mixing number of the VEGYTERV propeller has been investigated as a function of tank to agitator diameter ratio; Shiue and Wong [32] also found the same phenomena of 6-flat-blade turbine agitator. The mixing number depends on the low Reynolds number (Re < 100) of the reduced form [33] in the condition of gate type anchor impellers and multi-paddle agitators.

When the mixing time remains the same in the bioreactor at different scales, the P/V_L increases with two-thirds of the volume of the bioreactor [28]. Because high-volume power inputs are harmful to cells and do not require a constant mixing time to use as a scale up standard for mammalian cell culture of slow biologic reactions [30,31], which usually introduces concentration gradients forming and longer mixing times in large-scale bioreactors [31,32]. To improve the transport phenomenon, the dimensionless group is usually utilized in the mixing from small scale to large bioreactors [33,34]. The dimensional analysis principally presents that the dimensionless mixing number relies on Reynolds number and Froude number. On the other hand, bioreactors installed with baffles to escape vortex formation, and it is reasoned to be a function of Re as the free surface is nearly flat in laminar or transition flow systems [35]. However, for an orbital shaken reactor system acting with a non-flat free surface or with a vortex, Fr must also be considered as it describes the role of fluctuations on the free surface [36].

In stirred-tank bioreactor (STBs) cells, the high maximum power input value occurs at the stirrer tip and the bubble aeration foaming of the medium may result in high shear force. In comparison,
as a result of the power input distribution and the generalization of the bubble-free surface aeration, the shearing stress in OSBs is much lower. The mixing principle and the shear issue associated with STRs is well known to engineers [37–40], but results from studies with STBs cannot be applied to OSBs due to their agitating principle difference. Moreover, considering OSRs are relatively novel, the influence of crucial engineering parameters on bioreactor capability is yet to be well defined. This study aims to perform the mixing of Newtonian liquids in the turbulent regions of batched orbital shaken cylindrical bioreactors, describing the influences of various operating parameters on the power input and mixing time investigation at volumetric scales up to 20 L. Dimensionless Newton numbers and mixing number with regards to power consumption, mixing time, shaking frequency, bioreactor diameter, and fluid physical properties were utilized to determine power input and mixing time, and thereby mixing efficiency.

2. Materials and Methods

2.1. Equipment

The experiment was performed using a 20-L (28-cm-diameter × 51-cm height) cylindrical orbitally shaken bioreactor (Collect International Company) as described in Figure 1. The solutions utilized in this study include tap water and 5, 10 vol% and 20 vol% glycerin in water. All chemicals are industrial grade; Table 2 lists the viscosity and density of these solutions nearly Newtonian flow behavior. All measurements were performed at shaking frequencies of 80 rpm to 120 rpm, with filling volumes from 4 L to 10 L; shaking diameters of 3.2 cm, specifically adapted to mammalian cell culture processing.

![Figure 1. Schematic diagram of the experimental devices.](image)

| Liquid                  | Mimicking Culture Medium | Density * (kg/m³) | Viscosity * (Pa·s)(10⁻⁴) |
|-------------------------|--------------------------|-------------------|--------------------------|
| water                   | low 4% dextran 9000 [37] | 997               | 8.94                     |
| glycerin 5 volume%      | 0.5% polyvinylpyrrolidone 360 k [37] | 1022.75      | 13.53                    |
| glycerin 10 volume%     | 1.7% dextran 2,000,000 [37] | 1036              | 20.56                    |
| glycerin 20 volume%     | 2% dextran 2,000,000 [37] | 1051.95          | 23.87                    |

* Density is monitored using a Shimadzu TX223 L balance (the weight difference before and after divided by filling liquid volume), while viscosity is measured using a Physica MCR 301 rheometer.

2.2. Power Method for Power Consumption Determination

Experimental research on the local power consumption of the shaken bioreactor is based on the power functional method, with the specific steps during the experiment as follows:
a. The rotary shaking machine (rotating oscillator model: RTO-34) is firmly fixed to a frame [14]. The shaking table power input is regulated using a power meter (Arepu AR78-2, maximum loading power: 2200 W). The sealed vessel filled with a predetermined solution volume (4 L, 6 L, 8 L, 10 L) is placed on the rotary shaken platform. Measurements of the required power and speed data for shaking are recorded and saved on a personal computer [18];
b. Operate the closed vessel where the liquid moving is not permitted and then the electric power \( P_c \) is determined by multiplying the voltage and current values;
c. Operate the open vessel where the liquid can move freely and then the electric power \( P_o \) is calculated using the same method as step 2;
d. From the measured power consumption and the shaking frequency of the shaker drive (shaking diameter = 3.2 cm) local power consumption of \( \frac{P}{V_L} \) under this condition is determined [14, 38–41]:

\[
\frac{P}{V_L} = \frac{P_O - P_C}{V_L}
\]  

(1)

2.3. Temperature Method for Mixing Time Determination

A high temporal resolution, calibrated data logger with two external temperature probes (Commeter U0121) coupled with personal computer were used to measure the temperature difference and mixing time of two points within a bioreactor [25, 31–33, 42, 43]. To test the accuracy of this measurement technique, elementary experiments were carried out by setting two temperature probes in different locations as shown in Figure 1. The lower probe was placed at a distance of 0.12 m from the wall and 10 mm from the bottom of the reactor, while the upper temperature probe was located 10 mm below the liquid level at a distance of 0.14 m from the wall of the reactor. The probes location permitted an understanding of mixing dynamics within the reactor volume during mixing operation. The bridge temperature probes were utilized for high sensitivity to measure temperature differences approximate to 0.1 °C, and the temperature difference was recorded and calculated by the computer as a function of time. The quantitative data show there is no significant effect on the homogenization time of the turbulence area at the measuring point. Thermal pulse disturbance was used to the system by pouring into 2% of the same heated solution (60–70 °C) mixed at the midpoint of the liquid surface in the reactor. The temperature was controlled at 25 °C.

2.4. Theory

A dimensionless Newton number (\( Ne \)) related to the ratio of power resistance to inertial forces. It is utilized to represent the power input in the reactor. The Newton number of the power input is as follows [15]:

\[
Ne = \frac{P}{\rho \cdot N^3 \cdot D^4 \cdot V_L^{1/3}}
\]  

(2)

where \( P \) is power consumption, \( \rho \) is liquid density, \( N \) is shaking frequency, \( D \) is vessel diameter, \( V_L \) is filling volume.

The dimensionless Froude number (\( Fr \)) calculated using the ratio of inertial force to gravity. While the fluid is orbital moved in the shaken bioreactor, the number of free flow due to the centrifugal force of gravity acceleration must be modified to the radial direction ratio. The calculation of the axial Froude number is as follows [16]:

\[
Fr = \frac{2(\pi N)^2 \cdot D}{g}
\]  

(3)

where \( g \) is gravitational acceleration.

The dimensionless Reynolds number (\( Re \)) in fluid mechanics is a measurement of the ratio of inertial force to viscous force. The calculation of Reynolds number is as follows [15]:
\[ \text{Re} = \frac{\rho \cdot N \cdot D^2}{\eta} \]  

(4)

where \( \eta \) is dynamic viscosity of fluid.

The mixing number (Ho) is also known as the homogenization number [44], which is designated as the product of the mixing time and the shaking frequency and represents the shaking frequency required to achieve a given uniformity.

\[ Ho = N \cdot t_m \]  

(5)

where \( t_m \) is mixing time.

The correlation between volume power consumption and mixing time as follows [45]:

\[ t_m = C_1 \left( \frac{P}{V_L} \right)^c \]  

(6)

where \( C_1 \) and \( c \) are the constant and exponent, respectively.

The mixed effective number is the computational mixing efficiency of the reactor [46]:

\[ Ce = P_v \cdot t \]  

(7)

where \( P_v \) is the power consumption of unit volume, \( t \) is shaking time.

3. Results

Based on the number of variables to set up the new power input correlation for the Newton number (power number), the new mixing time correlation with volumetric number and volumetric power consumption and compare the mixing efficiency of different liquids and filling liquid volumes in the following relevant subsections. The intended correlation is a practical tool to figure the power input and mixing efficiency as cell cultivation in all OSB scales.

3.1. Power Input Correlation

3.1.1. Local Power Consumption

Power is transmitted during the shaking owing to the friction between the rotating liquid volume and the OSB wall. Different measurement systems used to determine the average power consumption in a OSB are introduced. A simple and efficient measurement of power consumption can be achieved by integrating a power meter in the OSB [15,18]. The method also permits online monitoring of power consumption during biological culture. However, integrating power meter into OSB is complex and requires a redesign of the OSB. The power consumption values in the cylindrical shaken reactor range from 37.1 W to 29.85 W, relying on the shaking frequency, filling volume and liquid viscosity. At 80 to 120 rpm shaking frequency, the filling volume for 20 L vessel influences local power consumption. Figure 2 shows local power consumption increases with the increased shaking frequency [14,15,41,47–50], but decreases with the increased filling volume [14,15,47,49] to be well mixed in-phase bulk liquid motion [49,51] as the friction surface between the OSB wall and the liquid is not proportional to the filling volume and consistent with the research of Büchs and Zoels [38] at shaking frequencies 50 to 400 rpm in 250-mL Erlenmeyer flasks in a 1 kg L\(^{-1}\) density organic dispersed phase of a mixture of four volumetric parts toluene and one part carbon tetrachloride. That power consumption significantly increases with viscosity in the case of 4 L filling volume is confirmed by the research of Büchs et al. [16] and Peter et al. [50] at the shaking frequency of 80 rpm to 300 rpm in a 300-mL baffled shake flasks using different viscosities polyvinylpyrrolidone (PVP) and shaking diameters of 100 mm. This is consistent with the research of Klöckner and Büchs [52] at shaking frequencies 80 to 300 rpm in 250-mL shake flasks, 40 mL filling volume and 25-mm shaking diameter in a viscous range of PVP. In case of 6 L to 10 L filling volume that the power consumption is
found to decrease with viscosity. Nevertheless, the shaking device is quickly accelerated, a low level of power consumption is reached away from a specific shaking frequency. Decreasing the shaking frequency below a critical value of 90 rpm suddenly produces an increase in power consumption reaching the standard level again [48]. The order of magnitude of the power consumption was shown with respect to shaking frequency and filling volume [53,54]. Power consumption exponentially increased with increasing shaking frequency [55]. The functional correlation between power consumption and shaking frequency is verified [14,51,56]

\[ P = A \cdot N^a \]  

where \( A \) is the coefficient, \( a \) is the exponent determined by experiments.

\( a \) is approximately 0.34 with a coefficient of determination greater than 0.9992 in a 20 L vessel with different fill volumes as shown in Figure 2.

3.1.2. Effect Reynolds Number on Newton Number

In the \( P/V_L \) correlation process, the number of variables is reduced by the theory of dimensionless analysis. We used the Buckingham \( \pi \)-theorem [57] to classify the variables as the following four dimensionless groups:

- Newton number: \( Ne = \frac{p}{\rho N^3 D^4 V_L^{1/3}} \)
- Reynolds number: \( Re = \frac{\rho N D^2}{\eta} \)
- Froude number: \( Fr = \frac{2(nN)^2 D}{g} \)
- Volume number: \( Vol = \frac{V_L}{D^3} \)
The power law is utilized as a mathematical correlation between the dimensionless number:

\[ Ne = B \cdot Re^\alpha \cdot Fr^\beta \cdot \left( \frac{VL}{D^3} \right)^\gamma \]  

(9)

where \( \alpha \), \( \beta \) and \( \gamma \) exponents are determined by experiments. One dimensionless number is modified in each experiment and other number remain the same.

The exponent in the Equation (8) must be determined by experiment. Thus, each dimensionless number is individually changed to distinguish the effects of each number in Equation (8). The correlation between Newton number and Reynolds number is first, evaluated, and the \( \alpha \) exponent of Equation (8) is determined. \( P/V_L \) is measured in 20 L vessel applying the liquids of different viscosity. The changes of liquid viscosity result in changes of Reynolds number without altering any other dimensionless number in Equation (8). Under different conditions, the \( \alpha \) value is verified using the shake frequency from 100 rpm to 120 rpm and the fill volume from 4 L to 10 L. The exponent \( \alpha \) of the Reynolds number is presented in Figure 3. As a result, the increase in Reynolds number corresponded to a decrease in Newton number \[18\]. The Newton number decreased with the increase in Reynolds number and Newton number of all liquids was the highest in 4 L, such as reported for an unbaffled mixing reactor \[58\]. The functional correlation between Newton number and Reynolds number is

\[ Ne = C_2 \cdot Re^\alpha = C_2 \cdot Re^{-2.66} \cdot R^2 = 1 \]  

(10)

where the ranges of Re is from \( 4.6 \times 10^4 \) to \( 1.75 \times 10^5 \) in the turbulent regime. This experiment results in a coefficient of determination greater than 0.9999.

![Figure 3. Relationship between Newton number and Reynolds number of different filling volumes. (a) Water; (b) 5 vol% glycerin; (c) 10 vol% glycerin; (d) 20 vol% glycerin.](image-url)
Equation (9) is confirmed by the research of Klöckner et al. [18] at shaking frequencies of 100 rpm, 120 rpm and 160 rpm in a 20-L vessel using polypropylene liquid with different viscosities and shaking diameters of 2.5 cm and 5 cm and consistent with the research of Meissner et al. [17] at shaking frequencies 160 to 340 rpm in 250 and 500-mL flasks in a viscous range of shear rates of polyvinylpyrrolidone fermentation broths between 0.1 s\(^{-1}\) and 3000 s\(^{-1}\). A variation of the liquid viscosity results in a variation of the Reynolds number without changing any other dimensionless number of Equation (9).

3.1.3. Effect Reynolds Number and Froude Number on Newton Number

For example, the different dual variables (Re and Fr) power correlations are researched with the minimum square regression by Klöckner et al. [18] at shaking frequencies of 100 rpm, 120 rpm and 160 rpm in a 20-L vessel using polypropylene liquid with different viscosities and shaking diameters of 2.5 cm and 5 cm and Scargiali et al. [59,60] report for the uncovered unbaffled and vortexing mixing reactor researched data, resulting in Equations (15) and (16), respectively.

\[
Ne = 2.4Re^{-0.17}Fr^{0.42} \quad (11)
\]

\[
Np = 24.0(Re \cdot Fr)^{-1/3} \quad (12)
\]

Therefore, all experimental Newton numbers are obtained versus the product (Re × Fr) and in very good agreement with Scargiali et al. [59,60] in the uncovered unbaffled and vortexing mixing reactor using \(10^{-3}\) to \(15.2 \times 10^{-3}\) Pa·s PVP and \(41.7 \times 10^{-3}\) to \(1100 \times 10^{-3}\) Pa·s glycerin solution at liquid height equal to tank diameter of 0.19 and 0.48 m. This experiment results in a coefficient of determination greater than 0.9999 as shown on Figure 4 and here can be represented Equation (13) (\(\alpha\) is equal \(\beta\) to \(-0.886\)), and a very good agreement can be observed.

\[
Ne = C_3(Re)^\alpha \cdot (Fr)^\beta \quad (13)
\]
3.1.4. Effect of Volume Number on Newton Number

The exponential $\gamma$ of the volume number is decided by changing the liquid volume $V_L$ [18]. Since merely the volume number is affected by the fill volume, exponential $\gamma$ is decided by changing the volume number only. Klöckner et al. [18] found that $Ne = 0.27 \cdot \left(\frac{V_L}{D^3}\right)^{0.44}$ was constant at shaking frequencies of 160 rpm, 20%–60% relative filling volume in a 20-L vessel using polypropylene liquid with different viscosities of 0.89–12 mPa·s and shaking diameters of 5 cm. The functional correlation between Newton number and volume number is obtained by

$$Ne = C_4 \left(\frac{V_L}{D^3}\right)^\gamma$$

(14)

The exponent of liquid volume in this correlation of the Newton number in our experiment differs from $-1.047$, $-1.1$, $-1.126$, $-1.205$ values with a coefficient of determination greater than 0.9999 obtained for different liquids as shown on Figure 5. This result is consistent with the investigation of the liquid distribution between shaking frequencies. Due to the cylindrical orbitally shaken bioreactor design, the liquid and reactor walls have higher liquid viscosity during shaking, so the volume number is more affected. Therefore, the Newton number as stated in Equation (2) is not suitable for OSB cylindrical vessels.

Figure 4. Relationship between Newton number and the product (Re x Fr) of different filling volumes. (a) Water; (b) 5 vol% glycerin; (c) 10 vol% glycerin; (d) 20 vol% glycerin.

Figure 5. Cont.
3.1.5. Validation and Application of Newton Number Models

The correlation of the power input in Equation (9) is established by experiment as following:

\[
Ne = 0.01117(Re \cdot Fr)^{-0.887} \left( \frac{V_L}{D^3} \right)^{-1.048} \text{ at water } (8.94 \times 10^{-4} \text{ Pa} \cdot \text{s})
\] (15)

\[
Ne = 0.0074(Re \cdot Fr)^{-0.887} \left( \frac{V_L}{D^3} \right)^{-1.116} \text{ at 5\% glycerin } (1.35 \times 10^{-3} \text{ Pa} \cdot \text{s})
\] (16)

\[
Ne = 0.0048(Re \cdot Fr)^{-0.887} \left( \frac{V_L}{D^3} \right)^{-1.14} \text{ at 10\% glycerin } (2.05 \times 10^{-3} \text{ Pa} \cdot \text{s})
\] (17)

\[
Ne = 0.004(Re \cdot Fr)^{-0.886} \left( \frac{V_L}{D^3} \right)^{-1.186} \text{ at 20\% glycerin } (2.4 \times 10^{-3} \text{ Pa} \cdot \text{s})
\] (18)

These equations were demonstrated to be valid in the following range:

\[
4.61 \times 10^4 < Re < 1.75 \times 10^5
\]

\[
1 < Fr < 2.3
\]

\[
0.18 < \frac{V_L}{D^3} < 0.46
\]

and when liquid motion was induced in the OSB that can be considered as being fully validated on scale-up.

3.2. Mixing Time

Mixing time was designated as the time beyond which at all measuring locations the thermal pulse disturbance differences between the temperatures measured and the final temperature was less than 3\% of the total temperature difference step (i.e., the difference between the initial and final temperatures) of the fluid. A typical example is expressed in Figure 6.
3.2.1. Effect of Shaking Frequency on Mixing Time

Two factors affecting the mixing time of the orbital shaking cylindrical bioreactor were studied. Because glycerin has a watery viscosity, it is expected that the mixing experiment in water is a well-being approximation of the mixing in the cell medium. Mixing time drops in log–log linearly with further increase at shaken frequency from 80 to 120 rpm [26,27,60]. Li et al. [24] determined that mixing time for $V_f = 6$-mL microwell plates was closely to 3.5 times greater than that of 2 mL at the lowest shaking frequency. Figure 7 shows the change in mixing time at shaking frequencies between 80 and 120 rpm at four different filling volumes (from 4 to 10 L), and the mixing time of $V_f = 10$ L being approximately 1.1-times the lowest shake frequency of 4 L at 20% high volume glycerin ratio.

![Figure 6. Typical example of the thermal pulse disturbance for the decision of mixing time. 20 vol% glycerin and 110 rpm shaking frequency.](image)

![Figure 7. Cont.](image)
3.2.2. Effect of Reynolds Number on Mixing Number

As shown in Figure 8, at the high Reynolds number of turbulent regime, the mixing number remains the independence of the Reynolds number nor dependent on proximity to the reactor wall since flow is turbulent [26,52]. Tan et al. [26] found that mixing number of the deionized water and viscous PVP solution were kept at approximately 20 and 10 at shaking frequencies of 100 to 350 rpm in a 100-, 250- and 500-mL standard unbaffled narrow-neck Erlenmeyer flasks using water and shaking diameters of 25 mm and 50 mm using 10%-filling nominal flask volume. The Reynolds number ranged at $10^4$–$10^5$ (turbulent region) and $10^2$–$10^3$ (turbulent to transition region), respectively. This evidence also shows that the mixing (homogeneous) number increases according to the liquid filling volume and the volume ratio of glycerin. Homogeneous number increased from 100.26 to 308.33, from 216.2 to 344.67, from 272.07 to 355.21 and from 298.47 to 366.17 for water, 5 vol% glycerin, 10 vol% glycerin and 20 vol% glycerin at four different filling volumes (from 4 to 10 L), respectively.

![Figure 7](https://example.com/figure7.png)

**Figure 7.** Effect of different shaking frequencies and filling volumes on mixing time. (a) Water; (b) 5 vol% glycerin; (c) 10 vol% glycerin; (d) 20 vol% glycerin.

![Figure 8](https://example.com/figure8.png)

**Figure 8.** Cont.
3.2.3. Effect of Volume Number on Mixing Time

It can be seen the mixing time for the bioreactor volume gradually increased when the ratio of filling volume to bioreactor volume increased from 0.18 to 0.46 [26]. The functional correlation between mixing time and volume number is obtained by

$$ t_m = C_5 \left( \frac{V_L}{D^3} \right)^b $$

(19)

where $V_L/D^3$ is volume number, $C_5$ is the coefficient, $b$ is the exponent is determined by experiments.

The exponent of volume number in this correlation of the mixing time in our experiment differs from 1.362, 0.509, 0.335, 0.21 values with a coefficient of determination greater than 0.9428 obtained for different liquids as shown on Figure 9. This result is consistent with the investigation of the liquid distribution between shaking frequencies. Due to the cylindrical orbitally shaken bioreactor design, the liquid and reactor walls have higher liquid viscosity during shaking, so the volume number is more affected.

Figure 9. Cont.
3.2.4. Effect of Volumetric Power Consumption on Mixing Time

The relationship between energy dissipation rate and mixing time was previously studied by Nienow [61]. To compare the mixing time of different liquids at the same power consumption, the relationship between mixing time and power consumption was plotted, results shown in Figure 10. With equal power consumption, the mixing time for water is the shortest. In addition, at lower power consumption, the mixing time increases according to liquid filling volume and the volume ratio of glycerin. This allows the prediction of the minimum mixing time for a given power consumption and demonstrates that more power consumption can improve the mixing process and reduce the mixing time in the reactor. A lower liquid viscosity was demonstrated to be more effective in liquid mixing given the shorter mixing time attained utilizing a lower power input at water, which is logical and desirable. In this experiment, the exponent of $P/V_L$ in the correlation of the mixing time is at $-2.93$ with a coefficient of determination greater than 0.9988 for water, 5 vol% glycerin, 10 vol% glycerin and 20 vol% glycerin on all shaking modes.
Figure 10. Effect of volumetric power consumption on mixing time. (a) Water; (b) 5 vol% glycerin; (c) 10 vol% glycerin; (d) 20 vol% glycerin.

There is symbolic difference between shake flasks and orbitally shaken bioreactor. Tan et al. [26] worked at shaking frequencies of 100 to 350 rpm in a 100-, 250- and 500-mL standard unbaffled narrow neck Erlenmeyer flasks using water and shaking diameters of 25 mm and 50 mm, concluding that $b = -0.58$ was constant. Therefore, in addition to the independence of the physical properties of the liquid, Equation (6) was found to be independent of liquid filling volume scale.

3.2.5. Validation and Application of Mixing Time Models

The correlation of the mixing time by combining the form of Equations (6) and (19) is established by experiment as following:

$$ t_m = 5.8685 \left( \frac{V_L}{D^3} \right)^{1.293} \left( \frac{P}{V_L} \right)^{0.3321} \quad \text{at water } \left( 8.94 \times 10^{-4} \text{ Pa} \cdot \text{s} \right) $$

$$ t_m = 3.1854 \left( \frac{V_L}{D^3} \right)^{0.482} \left( \frac{P}{V_L} \right)^{0.3059} \quad \text{at 5% glycerin } \left( 1.35 \times 10^{-3} \text{ Pa} \cdot \text{s} \right) $$

$$ t_m = 2.8893 \left( \frac{V_L}{D^3} \right)^{0.313} \left( \frac{P}{V_L} \right)^{0.2987} \quad \text{at 10% glycerin } \left( 2.05 \times 10^{-3} \text{ Pa} \cdot \text{s} \right) $$

$$ t_m = 2.766 \left( \frac{V_L}{D^3} \right)^{0.1805} \left( \frac{P}{V_L} \right)^{0.2808} \quad \text{at 20% glycerin } \left( 2.4 \times 10^{-3} \text{ Pa} \cdot \text{s} \right) $$

These equations were demonstrated to be valid in the following range and when liquid motion was induced in the orbitally shaken bioreactor:

$$ 4.61 \times 10^4 < \text{Re} < 1.75 \times 10^5 $$

$$ 1 < \text{Fr} < 2.3 $$

$$ 2.76 \text{ w m}^{-3} < \frac{P}{V_L} < 6.9 \text{ w m}^{-3} $$

$$ 0.18 < \frac{V_L}{D^3} < 0.46 $$

and when liquid motion was induced in the OSB that can be considered as being fully validated on scale-up.
3.3. Mixing Efficiency

In addition, mixing time can be measured and used to compare reactor efficiency at a given power consumption. The goal is to achieve the desired mixing strength with minimal power consumption. To compare the mixing efficient number of different filling liquid volumes, the relationship between the mixed effective number and the Reynolds number was plotted, as shown in Figure 11. It can be examined that the mixing effective number of water is the smallest in 4 L, while 5%, 10%, 20% vol. glycerin is the highest in 4 L, and the mixing effective number of all liquids is higher at the lower Reynolds number.

![Graphs showing mixing efficiency vs. Reynolds number for different liquid volumes.](a) Water; (b) 5 vol% glycerin; (c) 10 vol% glycerin; (d) 20 vol% glycerin.

4. Discussion

4.1. Effect of Operating Parameters on Newton Number

Local power consumption increases with increased shaking frequency but decreases with increased filling volume. The power consumption increases with viscosity in the case of 4 L filling volume, but in case of 6 L to 10 L filling volume, power consumption decreases with viscosity. Table 3 shows an overview concerning the functional correlation between power consumption and shaking frequency that was assessed in shaking flasks and orbital shaken reactors in previous studies. All experimental results in this study are in very good agreement with results presented in preceding works.
Table 3. Correlation between power consumption and shaking frequency.

| Bioreactor Type                      | Shaking Frequency | Shaking Diameter | Correlation | Liquid       | Literature               |
|--------------------------------------|-------------------|------------------|-------------|--------------|-------------------------|
| 20-L orbitally shaken bioreactor     | 100 rpm to 300 rpm| 50 mm            | P~N^{2.5}  | Tap water    | Kato et al. [48]        |
| 100- to 2000-mL flask                | 80 rpm to 380 rpm | 2.5 cm and 5 cm  | P~N^{2.8}  | Demineralized water | Büchs et al. [14]      |
| 20-L orbitally shaken bioreactor     | 300 rpm           | 2.5 cm           | P~N^{5.75} | Tap water    | Kato et al. [51]        |
| 20-L orbitally shaken bioreactor     | 80 to 120 rpm     | 3.2 cm           | P~N^{0.34} | 4.6 × 10⁴ to 1.75 × 10⁵ | This study              |

These authors respectively found that the functional correlation between Newton number and Reynolds number as presented in Table 4. Equation (9) was found to be independent of liquid filling volume. Büchs et al. [16] introduced the “in-phase” operating conditions for 100 to 2000-mL sizes of the flask and “out-of-phase” operating conditions symbolized by a non-developed liquid flow and irreproducibly occurred for 100- to 5-L size flasks [52], the correlation of modified Newton number and Reynolds number is

\[ Ne’ = 70 \cdot Re^{-1} + 25 \cdot Re^{-0.6} + 1.5 \cdot Re^{-0.2} \]  

(24)

Table 4. Correlation between Newton number and Reynolds number.

| Bioreactor Type                      | Shaking Frequency | Shaking Diameter | Correlation | Reynolds Number | Literature               |
|--------------------------------------|-------------------|------------------|-------------|-----------------|-------------------------|
| 100- to 2000-mL flask                | 80 to 380 min⁻¹ L/min | 2.5 and 5 cm | Ne’ = 70 Re⁻¹ + 25 Re⁻⁰.⁶ + 1.5 Re⁻⁰.² | 90 to 9 × 10⁴ | Büchs et al. [16]      |
| 250- and 500-mL flask                | 160 to 340 rpm    | 50 mm            | Ne = 12.3 Re⁻⁰.³⁶⁶       | 90 to 9 × 10⁴ | Meissner et al. [17]   |
| 20-L orbitally shaken bioreactor     | 100 rpm           | 2.5 cm           | Ne = 0.65 Re⁻¹.⁰.¹⁷     | 2.5 × 10⁴ to 3.5 × 10⁴ | Klöckner et al. [18]  |
|                                    | 120 rpm           | 2.5 cm           | Ne = 0.75 Re⁻¹.⁰.⁷      | 3 × 10⁴ to 5 × 10⁴    | Klöckner et al. [18]  |
|                                    | 160 rpm           | 5 cm             | Ne = 1.1 Re⁻¹.⁰.⁷      | 1.8 × 10⁴ to 3 × 10⁵  | Klöckner et al. [18]  |
| 20-L orbitally shaken bioreactor     | 80 to 120 rpm     | 3.2 cm           | Ne = C₁ Re⁻².⁶⁶        | 4.6 × 10⁴ to 1.75 × 10⁵ | This study             |

Equation (24) was utilized to consider in the laminar, intermediate and turbulent flow regime. Re⁻¹, Re⁻⁰.⁶ and Re⁻⁰.² terms in Equation (24) are related to the laminar, transitional and turbulent flow regimes, respectively.

Using Buckingham’s π theorem analysis, a new Reynolds number, the Froude number, and volume number are proposed for Newton number input in the cylinder orbitally shaken bioreactor. A new set of correlations have been proposed (Equation (15) to Equation (18)) for the Newton number in cylindrical orbitally shaken bioreactors. All experimental present Newton numbers are obtained versus the product (Re × Fr).

4.2. Effect of Operating Parameters on Mixing Time and Mixing Number

The mixing time is inversely proportional to the shaking frequency. At high Reynolds number, the homogenization number remains the constant. Tissot et al. [27] found that the mixing time remained to constant at higher shaking frequency in a 30-L vessel utilizing polypropylene liquid with different filling volume 8, 13 and 18 L. Mixing time of this experiment increases with the increased filling volume. Tan et al. [26] also found the mixing time for 100-, 250-, 500-mL nominal flask volumes increased gradually at 200 rpm shaking frequency. Consistently, the mixing in the viscous system, i.e., 20% high volume glycerin ratio solution at 10 L filling volume, was noticeably slower than in the non-viscous water system [26]. The mixing time increase phenomenon appears at a higher shaken frequency and with increasing fluid viscosity. For viscous fluids, the mixing time increase at high frequency range is also followed a decreased free surface vibration [24]. Under these conditions, the mixing number is independent of Reynolds number. This was also validated in traditional agitated tanks. The mixing (homogeneous) number increases according to increase the liquid filling volume.
and liquid viscosity. The mixing time for the bioreactor volume gradually increased as the ratio of filling volume to bioreactor volume is increased. The exponent of volume number and the mixing time correlation (Equation (19)) in the experiment obtained different values for different liquids. A new set of correlations were proposed (Equation (20) to Equation (23)) for the mixing time in cylindrical orbitally shaken bioreactors.

4.3. Effect of Operating Parameters on Mixing Efficiency

This study set out to compare the mixing efficiency of different liquids with the combination of power consumption and mixed time analysis. At lower power consumption, the mixing time increases according to liquid filling volume and liquid viscosity. This allows the prediction of the minimum mixing time for a given power consumption and demonstrates that more power consumption can improve the mixing process and reduce the mixing time in the bioreactor. Mixing efficient number shows that a constant power input of the reactor is fully consistent with the known trend of mixing time decrease.

5. Conclusions

Experimental studies of the power consumption and mixing time of the Newtonian fluid of water and glycerin in the cylindrical orbitally shaken bioreactor of the turbulent region were successfully completed. The power consumption and mixing time of continuous phase are determined by the power meter and the thermal method, respectively. The power consumption and mixing time are compared with the different volume of filled liquids. Using Buckingham’s π theorem analysis, a new Reynolds number and the Froude number are proposed for Newton number input in the cylinder orbitally shaken bioreactor. The mixing time is inversely proportional to the shaking frequency. At high Reynolds number, the homogenization number remains the constant. Under these conditions, the mixing time is independent of Reynolds number. The combination of power consumption and mixed time analysis of mixing efficient number shows that a constant power input, the mixer’s mixing time of the reactor is fully consistent with the known trend of the mixing time decreasing with the reactor size. Scale-up effects on the power consumption and mixing time in orbitally shaken bioreactor were investigated.

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Nomenclature

| Symbol | Description | Unit |
|--------|-------------|------|
| Ce     | computational mixing efficiency | W m⁻³ min⁻¹ |
| D      | vessel diameter | m |
| Fr     | Froude number | – |
| g      | gravitational acceleration | m s⁻² |
| N      | shaking frequency | rpm |
| Ne     | Newton number | – |
| P      | power consumption | W |
| Pᵥ     | power consumption of unit volume | W m⁻³ |
| Re     | Reynolds number | – |
| t      | shaking time | min |
| Vᴸ     | filling volume | m³ |

Greek Symbols

| Symbol | Description |
|--------|-------------|
| ρ      | liquid density | kg m⁻³ |
| η      | dynamic viscosity of fluid | Pa s |
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