Abstract—This paper reports on new approach of light illuminated esterification of oleic acid catalyzed by an enzymatic catalyst. Immobilized Pseudomonas cepacia lipase was used as a biocatalyst. Enzyme catalyzed esterification reaction was chosen due to the wide attention from chemical and biotechnology industry. The reaction was conducted under light illumination and dark condition at same experimental condition. The result was obtained in the form of changes in oleic acid concentration throughout 2h of reaction. The results indicate significant changes in rate of reaction for light illuminated reaction compare to reaction under dark condition. The initial rate of reaction for light induced and dark reaction are 1.2mol/L.min, 1.1mol/L.min respectively. Influence of enzyme amount was experimented where it shows proportional increase with the amount of enzyme. Increasing the amount of enzyme also increases the availability of active sites of enzyme therefore the probability of light illumination activating the electron transfer in oxyanion hole increases with increase in energy level of electron lone pair. The results obtained indicate that light illuminating method for enzyme activation provide good alternative for future biotechnology industry.

Index Terms—Esterification, light induced, oleic acid, Pseudomonas cepacia.

I. INTRODUCTION

Lipases are triacylglycerol acylhydrolases, E.C. 3.1.1.3 possess specific characteristics to catalyze ester hydrolysis reaction, esterification and transesterification reactions [1], [2]. These reactions widely explored by industries in recent years [3]. These are due to unique character of lipase structure being regiospecific, stereospecific and substrate specific [2], [4] and [5]. Furthermore, enzyme catalyzed reaction provides benefits over chemically synthesized reaction such as less energy condition by lowering the number of processes and it is environmental friendly [6].

However, due to the high enzyme production cost and low reaction rate the wide industrial application has not been implemented [7]-[9]. Therefore, researchers have been exploring the means to increase the rate of reaction by optimizing various parameters such as molar ratio, temperature, water content, immobilization method, pH and solvents for lipase from different sources.

However, none of the researchers implement light induced enzyme catalyzed esterification reaction until date. Hence, this study aims to activate the enzyme through light irradiation. Although, so far no studies on enzyme catalyzed esterification under visible light the novel approach was supported by previous researchers work on enzyme activation under the influence of light for different model. For instance, in 1986, Ljerka Purec et al. published that the hydrogenase activity of Proteus vulgaris could be enhanced by the irradiation of light with a wavelength of 253.7 nm. The studies also proposed that the light stimulation is irreversible reaction and the catalytic activity increases due to the conformational changes in enzyme [10], [11]. Later in 2003, Maciej Fiedorowicz et al. modified this light induced enzyme activity to study the effect of polarized and non-polarized visible light on hydrolysis of α-amylase. The authors showed that the conformational changes of α-amylase happened when the protein was illuminated with visible polarized and non-polarized light [12], [13].

Pseudomonas cepacia lipase was used in this study since its character as most adaptable biocatalyst. Furthermore, its crystal structure also reported by Kim et al. While P. glumae, R. miehei, G. candidum C. viscosum lipases show closed conformation where its active site structure is buried beneath a helical segment called as ‘lid’ and only opens up when interact with substrate [14]. Whereas, P. cepacia lipase illustrates highly open conformation made easier access of light to pass the active site of the enzyme.

Therefore, the objective of the study is to investigate the fundamental effect of light on enzymatic activity of lipases in esterification reactions. We have studied the lipase catalyzed esterification of oleic acid in the presence of commercially immobilized lipase from Pseudomonas cepacia in hexane under light and dark condition to identify the changes in rate of reaction for light illuminated enzyme catalyzed esterification reaction.

II. MATERIALS AND METHODS

A. Materials

P. cepacia lipase immobilized on chemically inert, hydrophobic sol–gel support was purchased from Sigma Aldrich (Switzerland). Ethanol, Oleic acid with an approximate purity of 99% and HPLC grade hexane were supplied by Fisher Scientific from Loughborough, United Kingdom. Sodium hydroxide was purchased from Merck KGaA, Germany. All chemicals, if not specified otherwise, were of analytical grade.

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B. Equipment and Esterification Reaction

Immobilized *P. cepacia* lipase was assayed in the direct esterification of oleic acid with ethanol to produce ethyl oleate and water as the end products.

Equation (1) describes the foremost esterification reaction between Oleic acid and ethanol with presence of lipase catalyst.

\[
\text{Lipase} \quad \text{Oleic acid} + \text{Ethanol} \rightarrow \text{Ethyl oleate} + \text{Water} \quad (1)
\]

The synthesis was carried out with 120mM oleic acid, 120mM ethanol and 0.11g *Pseudomonas Cepacia* lipase in isothermal batch type reactor with continuous stirring at 300rpm equipped with a heat transfer jacket and cooling system as shown in Fig. 2. All the reaction was conducted at 37°C. In a typical experiment, the reactants were introduced in the reactor and heated up to the working temperature. Once the reaction temperature was reached, the reaction was activated by adding the immobilized *Pseudomonas cepacia* enzyme into the reactor.

C. Analysis of Reaction Product

The progress of the esterification was monitored by determination of the remaining oleic acid content by titrating with a 0.01M basic solution of sodium hydroxide. Phenolphthalein was used as the end-point indicator and a mixture of ethanol-acetone 50/50% (v/v) was used as quenching agent. When the enzymatic activity stopped, the catalyst was separated by filtration, washed with ethanol (96%) and dried. All the experiments were repeated twice.

III. RESULTS AND DISCUSSION

A. Synthesis of Ethyl Oleate

Fig. 3 shows the reduction in concentration of oleic acid with respect to time. The reaction was conducted under both visible light and dark condition with total reaction volume of 12ml hexane consist of 120mM oleic acid, 120mM ethanol with presence of 0.11g immobilized *Pseudomonas cepacia* enzyme. As shown in Fig. 1. After 2 h of reaction more than 50% conversion was obtained for light and dark reactions. However, there is significant change in concentration of oleic acid presence in reaction volume for the first 20min in light illuminated experiment. This may be due to acceleration of oleic acid transferring to histidine residue in oxyanion hole with light illumination. The crystal structure of *P. cepacia* lipase reported by Kim *et al.* illustrates open conformation structure (open lid) which made easy reachable of light on active site.

The rate was calculated using the three point differentiation formula as shown below:

\[
\left( \frac{dCA}{dt} \right)_{t_o} = \frac{-3CA_0 + 4CA_1 - CA_2}{2\Delta t}
\]

The value of Table I was extracted from Fig. 2. Using (1) where light has high initial rate compare to dark reaction with 1.200 mol/L.min and 1.113 mol/L.min.
B. Influence of Enzyme Amount

The significant increase for light induced reaction could be due to excitation of electron lone pair on histidine-nitrogen during tetrahedral intermediate of oxyanion hole which could move to higher energy level resulting into facilitated passing through transition state of the process for formation of a covalent bond between histidine-nitrogen and hydrogen from serine. Therefore, the excitation of electron enhances the activity of lipase enzyme to accelerate the reaction rate [14].

IV. CONCLUSION

The objective of this study is to evaluate on rate of esterification reaction under light illumination. It was demonstrated that light induced reaction has higher rate of reaction compared to the dark reaction. Hereafter, this method could be implemented for enzyme catalyzed hydrolysis and transesterification reaction. Unambiguously, light illumination for lipase catalyzed reactions should take into serious consideration for future of lipase enzyme catalysis especially for biofuel production.

REFERENCES


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