PARTIAL PURIFICATION AND CHARACTERIZATION OF MOLYBDENUM REDUCING ENZYME FROM KLEBSIELLA OXYTOCA STRAIN HKEEM

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FBSB 2011 20
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MASTER OF SCIENCE
UNIVERSITI PUTRA MALAYSIA

October 2011
Dedicated to my parents, family and friends
Abstract of thesis presented to the Senate of Universiti Putra Malaysia in fulfilment of the requirements for the degree of Master of Science

PARTIAL PURIFICATION AND CHARACTERIZATION OF MOLYBDENUM REDUCING ENZYME FROM KLEBSIELLA OXYTOCA STRAIN HKEEM

By

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October 2011

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As a result of widespread application in numerous industrial processes, heavy metals have become a contaminant of many environmental systems. Molybdenum’s ubiquitous application in many industrial products makes it a silent pollutant with levels as high as several hundreds of ppm. Thus in this study, the isolate molybdenum reducing bacterium, which showed the highest molybdenum reducing activity, were isolated from a soil sample collected near a steel factory in Selangor. This isolate was later identified as Klebsiella oxytoca strain hkeem through molecular phylogenetic analysis of sequenced 16S rRNA gene sequence. Klebsiella oxytoca strain hkeem reduced the heavy metal molybdenum to molybdenum blue optimally in selective low phosphate medium agar, at pH 7.3 and 30°C, after 24 hours of incubation. In specific, Klebsiella oxytoca strain hkeem could reduce molybdenum to molybdenum blue under aerobic conditions in the medium with fructose as electron donor, yeast extract, phosphate ion (4.5 mM) and molybdate ion (80 mM). Partial purification and characterization were conducted on the molybdenum reducing enzyme, with anion exchange on Q-sepharose and gel filtration on Zorbax GFX-250. Based on the SDS polyacrylamide-gel
electrophoresis (SDS-PAGE), two bands were observed on the gel filtration fraction at 90 and 38 kDa, respectively. Meanwhile, the enzyme showed an optimum activity at substrate pH 5.5 and 25ºC. The kinetics of electron donor (NADH) and electron acceptor (LPPM) for the enzyme were followed the classical Michaelis-Menten rectangular hyperbolic curve. $K_m$ and $V_{max}$ for the electron donor substrate, NADH was 2.83 mM and 12.23 nmole molybdenum blue produced/min/mg/protein, respectively. However the $K_m$ and $V_{max}$ for the electron acceptor substrate phosphomolybdate (50 mM) were 1.66 mM and 32.06 nmole molybdenum blue/min/mg/protein respectively. Although many molybdenum-reducing bacteria have been isolated, molybdenum-reducing activity of *Klebsiella oxytoca* strain hkeem up to 80 mM of molybdate was reported as the most potent molybdenum-reducing isolate to date. Based on the results obtained, *Klebsiella oxytoca* strain hkeem was proven as a more powerful molybdenum reducer to develop a cost-effective bioremediation work, especially since bacterial molybdenum reduction has been suggested as an important remediation tool for cleaning up molybdenum pollutant in the environment.
PENULENAN SEPARA DAN PENCIRIAN ENZIM PENURUM MOLYBDENUM DARIPADA STRAIN KLEBSIELLA OXYTOCA HKEEM

Oleh

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Oktober 2011

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Akibat daripada penggunaan yang berleluasa dalam pelbagai proses perindustrian, logam berat telah menjadi bahan pencemar alam sekitar yang utama. Penggunaan molybdenum yang meluas dalam pelbagai produk industri, menjadikan molybdenum bahan pencemaran alam sekitar. Oleh itu, dalam kajian ini bakteria penurun molybdenum yang menunjukkan aktiviti penurunan molybdenum yang tinggi telah dipencilkan daripada sampel tanah berdekatan dengan kilang besi di Selangor. Pencilan ini dikenalpasti sebagai strain hkeem Klebsiella oxytoca melalui filogenetik molekul menggunakan jujukan gen 16S rDNA. Klebsiella oxytoca strain hkeem menurunkan logam berat molybdenum ke molybdenum biru dengan optimum pada agar media terpilih yang rendah fosfat, pada pH 7.3 dan 30ºC, selepas pengeraman selama 24 jam. Secara khasnya, Klebsiella oxytoca strain hkeem boleh menurunkan molybdenum ke molybdenum biru dibawah keadaan aerobik dalam media yang mengandungi fruktosa sebagai penderma elektron, ektrak yis, cas fosfate (4.5 mM) dan cas molybdate (80 mM). Penulenan separa dan pencirian enzim penurun molybdenum telah dijalankan dengan menggunakan penukar anion pada Q-sepharose dan penapisan gel pada Zorbax GFX-250. Berdasarkan elektroforesis SDS polyacrylamide-gel (SDS-PAGE), 2 jalur...
diperhatikan pada bahagian penapisan gel pada 90 dan 38 kDa. Sementara itu enzim ini menunjukkan aktiviti tertinggi substrat ialah pada pH 5.5 dan 25°C. Penderma elektron (NADH) dan penerima elektron (LPPM) enzim tersebut adalah berdasarkan kepada lengkungan hiperbolik segiempat Michaelis-Menten klasik. $K_m$ dan $V_{max}$ untuk substrat penderma elektron ialah, NADH ialah 2.83 mM dan 12.23 nmole molybdenum biru terhasil/min/mg/protein. Manakala $K_m$ dan $V_{max}$ penerima elektron fosfomolybdat (50 mM) ialah 1.66 mM dan 32.06 nmole molybdenum biru/min/mg/protein. Walaupun banyak bakteria penurun molybdenum telah dipencarkan sebelum ini, keupayaan *Klebsiella oxytoca* strain hkeem menurunkan molybdenum sehingga 80 mM molybdat merupakan bakteria penurun molybdenum yang paling poten dipencarkan setakat ini. Bio-pemulihan telah dicadangkan sebagai kaedah yang lebih kos efektif dan penting untuk membersihkan bahan pencemar alam sekitar. Berdasarkan keputusan yang diperolehi dalam kajian ini, *Klebsiella oxytoca* strain hkeem telah terbuktikan sebagai penurun molybdenum yang paling berpotensi untuk membangunkan teknologi bio-pemulihan ini.
ACKNOWLEDGEMENTS

Writing the acknowledgement is a wonderful phase to express in so few words to all the deepest appreciation people who made this Msc. Dissertation completed. I owed my greatest thanks, appreciation and admiration to my family. They have given me their endless support, motivation and love not only during the completion of this thesis, but throughout my lifetime. I’m sure they will continue showering me with their love as I pursuing all my dreams in my life. They inspired me to be a better person.

I would like to convey my deepest gratitude and appreciation to my main supervisor, Assoc. Prof. Dr. Mohd Yunus Abd. Shukor and also not forgotten my Co-supervisor Prof. Dr. Mohd Arif Syed. Their advise, priceless guidance and patience that they have given me throughout the entire duration has make me becoming more passionate and dedicated to finish this thesis. Again, thank you so much.

I also want to thank my fellow friends in Lab 204 (Bioremediation Lab) and Lab 115 (Enzymology Lab) for supporting and helping me in finishing my lab works. Special thanks to the members of molybdenum group member. They have helped a lot by sharing their knowledge, ideas and experiences. They also helped me to overcome obstacles and difficulties that I had gone through with this thesis. They are not only research members but friends for a lifetime. Special thanks to my mentor of my life Mr. Daisaku Ikeda. He is the philosopher. His encouragement in his writing had encouraged me not only when I facing the problems in this master study but also in my life.
I certify that an Examination Committee has met on 14th October 2011 to conduct the final examination of Lim Hui Keem on her thesis entitled “Partial Purification and Characterization of Molybdenum-reducing enzyme from *Klebsiella oxytoca* strain hkeem” in accordance with Universities and University Colleges Act 1971 and the Constitution of the Universiti Putra Malaysia [P.U.(A) 106] 15 March 1998. The Committee recommends that the student be awarded the Master of Science degree.

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DECLARATION

I declare that the thesis is my original work except for quotations and citation which have been duly acknowledged. I also declare that it has not been previously and is not currently submitted for any other degree at Universiti Putra Malaysia or other institutions.

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LIM HUI KEEM
Date: 14 October 2011
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