

FACULTY OF RESOURCE SCIENCE & TECHNOLOGY





Research in biotechnology is at the forefront of scientific innovation, combining biology, chemistry, and engineering principles to develop revolutionary solutions that impact various sectors, from healthcare to agriculture and environmental conservation. At Faculty of Resource Science & Technology, UNIMAS, we are dedicated to advancing knowledge and pushing the boundaries of this exciting field.

WHY BIOTECHNOLOGY RESEARCH IS IMPORTANT?

Advancing Healthcare

Biotechnology plays a vital role in developing new therapies, diagnostics, and vaccines. Through genetic engineering and biopharmaceutical production, we are paving the way for personalized medicine, targeted treatments, and more effective disease management.

Sustainable Agriculture

Biotechnology enables the development of genetically modified crops that exhibit enhanced yield, resistance to pests, and tolerance to environmental stress. This has the potential to address food security challenges, reduce reliance on chemical pesticides, and promote sustainable agricultural practices.

Environmental Conservation

Our research focuses on leveraging biotechnology to address environmental issues. From bioremediation techniques that aid in cleaning up polluted sites to biofuels and renewable energy production, we strive to minimize the impact of human activities on the planet.

Industrial Applications

Biotechnology research has a impact on profound various industries. From the production of bio-based materials and biofuels to the development of novel enzymes biocatalysts for industrial and processes, our work contributes to a more sustainable and efficient industrial landscape.





























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NICHE: MICROBIOLOGY

SOME OF OUR RECENT WORKS

Research Article

Prevalence, Genetic Heterogeneity, and Antibiotic Resistance Profile of Listeria spp. and Listeria monocytogenes at Farm Level: A Highlight of **ERIC- and BOX-PCR to Reveal Genetic Diversity**

Lesley Maurice Bilung ,1 Lai Sin Chai,1 Ahmad Syatir Tahar,1 Chong Kian Ted 0,2 and Kasing Apun 0

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circu. This study aimed to identify *Listeria* spp. and *L. monocytogenes*, characterize the isolates, and determine the profiles of the isolates *Listeria* spp. and *L. monocytogenes* in fresh produce, fertilizer, and environmental san farms (organic and conventional farm). A total of 386 samples (vegetables, soil, water, and fertilizer with manure) were examined. The identification of bacterial isolates was performed using PCR and characterized using ERIC-PCR and BOX-PCR. The discriminating power of the typing method was analyzed using Simpson's Index of Diversity. If were subjected to antimicrobial susceptibility test using the disc-diffusion technique. The PCR a were present in 251% (29/386) of all the samples (vegetable, soil, fertilizer, and water). None of the for the presence of *L. monocytogenes*. Precentages of 100% (151%) and 73.30% (11/5) of the *Lister* fertilizer, and soil from organic farm B had indistinguishable DNA fingerprints by using ERIC-1 *Listeria* spp. isolated from 86 samples of vegetable, of tilizer, and environment of organic farm distinct DNA fingerprints. Simpson's Index of Diversity, D. of ERIC-PCR and BOX-PCR is 0.6044 susceptibility test revealed that most of the *Listeria* spp. in this study were found to be resistant to interesting the distingtion of the *Listeria* spp. The subject of the subject of the *Listeria* spp. The subject of the subje

INTERNATIONAL JOURNAL OF BIOLOGY AND BIOMEDICAL ENGINEERING DOI: 10.46300/91011.2022.16.37

Volume 16, 2022

Contamination of waterborne parasites at water treatment plants and a gravity-feed system: a highlight on water safety for urban and rural communities in Kuching, Sarawak

Ahmad Syatir Tahar¹, Lesley Maurice Bilung¹, Kasing Apun¹, Reena Leeba Richard³, Hashimatul Fatma Hashim¹, Elexson Nillian¹, Lau Seng⁴, Yvonne Ai-Lian Lim²

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International Food Research Journal 24(3): 985-989 (June 2017)

Detection of Bacillus cereus in formula milk and ultra high temperature (UHT) treated milk products

1*Lesley, M. B., ¹Ernie, S.R., ¹Kasing, A. and ^{2,3}Son, R.

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Article history

Abstract

ed: 20 May 2016 ed in revised form ne 2016 pted: 9 June 2016

Keywords

Bacillus cerens Infant formula

Ultra high temperature (UHT) treated milk products and formula milk are known to be frequently contaminated with Bacillus cereus. Presence of B. cereus in these milk products is requently containing of the second se at detecting gyrB gene in B. cereus, that encode for the subunit B protein of DNA gyrase (tensiscentration to B). Contemination load of B contemination load of B.

of Advanced Research in Applied Sciences and Engineering Technology 11, Issue 1 (2018) 47-5-



Growth Kinetics of Ethidium Bromide Mutagenized Lipomyces starkevi Strains

Micky Vincent^{1,*}, Latifah Suali¹, Afizul Safwan Azahari¹, Patricia Rowena Mark Baran¹, Elexson Nillian¹, Lesley Maurice Bilung¹

nt of Molecular Biology, Faculty of Resource Science and Technology, Universiti Malaysia Sarawak, 94300 Kota Samarahan, Sarawak,

ARTICLE INFO	ABSTRACT
Article history: Beceived 17 January 2018 Beceived In revised from 28 February 2018 Accepted 2 May 2018 Available online 13 May 2018	Yeast growth and biomass production are greatly influenced by the length of the incubation period during cultivation. Therefore, this study was conducted to investigate the growth kinetics of five <i>Lipomyces storkeyi</i> strains as determined by biomass production. The five <i>L.</i> storkeyi artise, namely <i>L.</i> storkeyi ATCC 12659, <i>L.</i> storkeyi MV-1, <i>L.</i> storkeyi MV-4, <i>L.</i> storkeyi MV-5 and <i>L.</i> storkeyi MV-8, were inoculated in sterilized Yeast Mali borch, and, incubated for 132 hr at ambient temperature. Biomass yields were assessed and calculated gravmetrically every 24 hr. Results indicated that the noticeab fiberary storketing of <i>L.</i> other <i>MCC</i> 13569. <i>L.</i> torketing

High Occurrence of Staphylococcus aureus Isolated from Fitness **Equipment from Selected Gymnasiums**

> ley Maurice Bilung 💿, Ahmad Syatir Tahar, Rosdi Kira, Aina Ariffah Mohd Rozali, Kasing Apun

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ived 29 March 2018; Revised 17 July 2018; Accepted 2 August 2018; Published 28 August 2018

lemic Editor: Giuseppe La Torre

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sduction. Staphylococcus aureus is a leading cause of cutaneous bacterial infection involving community. Methods. In this y, a total of 42 swab samples were collected from the surface of various fitness equipment such as back machines, exercise , dip stations, dumbbells, and treadmills. Identification of the bacterial isolates was conducted using biochemical tests and er analysed molecularly using the PCR method targeting nuc gene (270 bp). The nuc gene encodes for the thermonuclease me, a virulent factor of S. aureus. Results. The findings showed 31 out of 42 swab samples (73.81%) were positive with resu. Conclusion. This study showed that gymnasium equipment is a potential reservoir for S. aureus and might play-ah/V/ir

BIODIVERSITAS ser 4, July 2018 Volume 19, Numb Pages: 1266-1272

ISSN: 1412-033X E-ISSN: 2085-4722 DOI: 10.13057/biodiv/d190412

Isolation, identification and diversity of oleaginous yeasts from Kuching, Sarawak, Malaysia

MICKY VINCENT", HUANG CHAI HUNG, PATRICIA ROWENA MARK BARAN, Y VINCENT, HUANG CHARIFORD AND SALWANI AWANG ADENI AFIZUL SAFWAN AZAHARI, DAYANG SALWANI AWANG ADENI Adapting Ditermit Malawin Sarawak, 94300 Ko arce Bioteche an. Sarawak

> Mari script received: 25 May 2018. Revision accepted: 6 June 2018

Ubstract. Vincent M, Hung MC, Baran PRM, Azahari AS, Adeni DSA. 2018. Isolation, identification and diversity of oleaginous yeasts rom Kuching. Sarawak. Malaysia. Biodiversitas 19: 1266-1272. The present study was performed to isolate, identify and determine the rom Kuching. Sarawak.

Pusigraduate Symposium in Busechnology (IPSB) 2019	gth, typical of
XA AND CTXAB) GENES IN ROM CLINICAL AND	yeast isolates, (21), Candida es marxianus n IV staining.
MBANG SARAWAK BY	

Amirah, Z. J.¹, Elexson, N.^{1*}, Grace, B.¹, Diyana, Z.¹

MULTIPLEX POLYMERASE CHAIN REACTION (PCR)

«Proceeding 3nd Inte

DETECTION OF CHOLERA TOXIN (CT) **VIBRIO CHOLERAE ISOLATED F** ENVIRONMENTAL SAMPLES IN LI

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ABSTRACT

ADSTRACT Cholera epidemics have been occurred in Malaysia since 1991 till 2003 which can be proved from the records by the Infectious Diseases Division of the Ministry of Health. Moreover, there were also course of cholera epidemics from the year 1994 to 2003 had been described in Sarawak. Cholera outbreaks in Malaysia mostly occur caused by the El Tor OI. *Vibrio cholerae* serogroup. Previous research has been focusing on the presence of violations and to detect associated factors in *V. cholerae* populations include cholera toxin (czrA). The aims of this study were to detect the mesence of *Vibrio cholerae* in clinical and environmental samples (n=28) from Limbang Sarawak and to detect

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Biotechnology plays a vital role in ensuring food security by increasing crop yields, improving crop quality and nutritional value, enhancing pest and disease resistance, promoting sustainable agriculture practices, and conserving genetic resources. These advancements contribute to a more resilient and productive agricultural system, helping to meet the global demand for safe, nutritious, and affordable food while addressing the challenges posed by population growth, climate change, and limited arable land.

NICHE: FOOD SECURITY

SOME OF OUR RECENT WORKS



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UPM Serdang, Malaysia

Graphical abstract Abstract



CR product sizes of mitochondrial DNA cvtochrome b gene for the investigated Through the advancement of effective techniques in specie stability in harsh conditions co conducted to determine the e phenol/chloroform/isaamyl a commercially available kit by for amplification of porcine DI and quality of the DNA extract Polymerase chain reaction (P targeting mitochondrial DNA co DNA extracts. High DNA con samples extracted using the f samples extracted using the l agarose gel was able to dete minute proportion (1%). The e ith the presence of a spe recommended as a cost-effect extraction kits in detecting port

Keywords: Meat adulteration cytochrome b

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Detection of Beneficial Lactic Acid Bacteria (LAB) and Yeast In Sarawak Fermented Food

Elexson Nillian1,*, Nick Laurence Buyong1, Dalene Lesen1, Grace Bebey1, Azham Zulkharnain2

- Faculty of Resource Science and Technology, University Malaysia Sarawak, 94300, Kota Samarahan, Sarawak, Malaysia
- ² Department of Bioscience and Engineering, Shibura institute of Technology, Japan

ABSTRACT

Sarawak native's fermented food can be a catalyst for boosting the local economy in Sarawak. The Lactic Acid Bacteria (LAB) are generally regarded as safe, have a stability of usage, and originate from natural resources. Lactic acid bacteria and yeast work in synergy to provide a natural way to enhance the nutritive value and flavour of the food. The study aims to investigate the presence of potential probiotic Lactic Acid Bacteria (LAB) and yeast isolated from Sarawak fermented food. Two hundred fifty (n=250) of ding fifty (n=50) each sample as fe nted shrin ns (cencaluk) fermented musta etables (kas



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Plant biotechnology is essential for sustainable agriculture, improved crop productivity, enhanced nutrition, environmental conservation, and adaptation to a changing climate. It offers innovative solutions to address global food security challenges, reduce environmental impacts, and promote the sustainable use of plant resources for the benefit of both human welfare and the planet.

NICHE: PLANT BIOTECHNOLOGY

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< PLANT BIOLOGY

In vitro induction and identification of polyploid *Neolamarckia cadamba* plants by colchicine treatment

arch article Agricultural Science Biotechnology Plant Science

Wee Hiang Eng¹, Wei Seng Ho^{™1}, Kwong Hung Ling²

Published October 27, 2021

> Author and article information

Abstract

Polyploidization has played a crucial role in plant breeding and crop improvement. However, studies on the polyploidization of tropical tree species are still very scarce in this region. This paper described the *in vitro* induction and identification of polyploid plants of *Neolamarckia cadamba* by colchicine treatment. *N. cadamba* belongs to the Rubiaceae family is a natural tetraploid plant with 44 chromosomes (2n = 4x = 44). Nodal segments were treated with colchicine (0.1%, 0.3% and 0.5%) for 24 h and 48 h before transferring to shoot regeneration medium. Flow cytometry (FCM) and chromosome count were employed to determine the ploidy level and chromosome number of the regenerants, respectively. Of 180 colchicine-treated nodal segments, 39, 14 and 22 were tetraploids, mixoploids and octoploids, respectively. The highest percentage of



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Home > The Protein Journal > Article

Published: 20 December 2019

Proteomics of Sago Palm Towards Identifying Contributory Proteins in Stress-Tolerant Cultivar

Hasnain Hussain ⊠, Maswida Mustafa Kamal, Jameel R. Al-Obaidi, Nur Ezzati Hamdin, Zainab Ngaini & Yusmin Mohd-Yusuf

The Protein Journal 39, 62-72 (2020) Cite this article

390 Accesses | 6 Citations | Metrics

Abstract

Metroxylon sagu Rottb. or locally known as sago palm is a tropical starch crop grown for starch production in commercial plantations in Malaysia, especially in Sarawak, East Malaysia. This plant species accumulate the highest amount of edible starch compared to other starch-producing crops. However, the non-trunking phenomenon has been observed to be one of the major issues restricting the yield of sago palm starch. In this study, proteomics approach was utilised to discover differences between trunking and non-trunking proteomes

Scientia Horticulturae 246 (2019) 604-617

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17-18 ·	Scientia Horticulturae	
ELSEVIER	journal homepage: www.elsevier.com/locate/scihorti	

Polyploidization using colchicine in horticultural plants: A review

ABSTRACT

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A R T I C L E I N F O Krywinfie Cheensoone doubling Antinistoite agent Polyploid Flow cytometry Cheensownie counting

Polypholdization of plants occurs in nature but it is a dow and gradual process that has driven evolution and speciation throughout the ages. At time is important, polypholdization can be induced in a thereter period by using colchicities that interferes with mitosis of cells. To develop an efficient polypiolitation portcoid of a species, up to date research is parameters. The efficiency of polypioldization induction system depends on the advancement of progragation techniques of the species which consist of invitor, or wire and in the systems. During polypiolitation induction, several parameters can be inneipialed to obtain highest percentage of ophyladis. These parameters include colchicien concentration, transmeter duration galaxie types. Several ophyladis distributes of several parameters in schedule collicities concentration, transmeters, aniotanical parameters and flow cylinodization has been practical for decades with notable breakthough recently using chromosome consting. Polypiolidization has been practical for decades with notable breakthough recently using chromosome consting. Polypiolidization has been practical for decades with notable breakthough recently in terms of polypiolidization system and polypiolid disattication. Current trend in polypiolidization has included to polypiolidized breakthough recently in a several polypiolidization has included to polypiolidized breakthough recently in the polypiolidization brack hough polypiolidization terms of polypiolidization system and polypiolidization for decades with notable breakthough recently in terms of polypiolidization present and polypiolidization and the polypiolidization terms of polypiolidization polypiolidization terms of polypiolidization polypiolidization polypiolidization polypiolidization polypiolidization terms of polypiolidization polypiolidiza

Nisar and Hussain Chem. Biol. Technol. Agric. (2022) 9:46 https://doi.org/10.1186/s40538-022-00315-1

Chemical and Biological Technologies in Agriculture

Open Access

RESEARCH

Assessment of the genetic variations of sago palm *Metroxylon sagu* in three regions of Sarawak, Malaysia using amplified fragment length polymorphism (AFLP) marker

Mehvish Nisar[®] and Hasnain Hussain[®]

Abstract

Background: Sago palm (Metroxylon sogu) is an important staple crop in the rural area of Sarawak, Malaysia. The palms grow well in the rough, swampy peat, from low flooded areas to uplands and acidic to neutral soils. Among the starch-producing crops, this palm is the most productive and promising, where it can store a significant amount of carbohydrate in the trunk. Due to the lack of molecular study of sago palm, the work described here aimed to develop molecular markers for identifying *M. sagu* and accessing the genetic variations of sago palm in different locations of Sarawak.

Results: Amplified fragment length polymorphism was employed to determine the genetic variations of sago

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Biotechnology offers novel resources to enhance the well-being of humans and animals, as well as to boost livestock productivity. It contributes to the improvement of our food sources, such as meat, milk, and eggs. Additionally, biotechnology can positively influence an animal's environmental footprint. Moreover, it enhances our capacity to identify, treat, and prevent diseases.



NICHE: ANIMAL BIOTECHNOLOGY

SOME OF OUR RECENT WORKS



A Review on the Emerging Asian Aquaculture Fish, the Malaysian Mahseer (Tor tambroides): Current Status and the Way Forward

Melinda Mei Lin Lau 🖾 Leonard Whye Kit Lim 🖾 Sairatul Dahlianis Ishak, Ambok Bolong Abol-Munafi & Hung Hui Chung

ceedings of the Zoological Society, 74, 227–237 (2021) Cite this article 670 Accesses | 9 Citations | Metrics



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Gene Reports Volume 23, June 2021, 101110



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Zebrafish (Danio rerio) ecotoxicological ABCB4, ABCC1 and ABCG2a gene promoters depict spatiotemporal xenobiotic multidrug resistance properties against environmental pollutants

Leonard Whye Kit Lim⁴ & 🗃 , Hung Hui Chung⁴ & 🗃 , Sairatul Dahlianis Ishak⁶ , Khor Waiho⁶ Show more V



https://doi.org/10.1016/j.genrep.2021.101110 >

Abstract

Marine organisms are naturally equipped with multixenobiotic resistance mechanisms that are often governed by ATP-binding cassette (ABC) transporter family members. Previous studies focused on the target genes of ABC but little is known about the functionality of their promoter regions. Due to the importance of promoters in ABC

First high-quality genome assembly data of sago palm (Metroxylon sagu Rottboll)

Leonard Whye Kit Lim^a 2 🖾 🖶 , Melinda Mei Lin Lau^a, Hung Hui Chung^a 2 🖾 , Hasnain Hussain^b, Han Ming Gan^{cd}

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https://doi.org/10.1016/j.dib.2022.107800 >



Data in Brief Volume 39, December 2021, 107481



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Data Article

The first transcriptome sequencing and data analysis of the Javan mahseer (Tor tambra)

Melinda Mei Lin Lau³, Leonard Whye Kit Lim³, Hung Hui Chung³ 🙎 🖾 , Han Ming Gan^{b c}

- 10				
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Abstract

The Javan mahseer (Tor tambra) is one of the most valuable freshwater fish found in Tor species. To date, other than mitogenomic data (BioProject: PRJNA422829), genomic and transcriptomic resources for this species are still lacking which is crucial to understand the molecular mechanisms associated with important traits such as growth, immune response, reproduction and sex determination. For the first time, we sequenced the transcriptome from a whole juvenile fish using Illumina NovaSEQ6000 generating raw paired-end reads. De novo transcriptome assembly generated a draft transcriptome (PLKCOE completeness of 01.2% [Actingstappeli adb10 database]) consisting of 250.402

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SOME OF OUR RECENT WORKS

Jurnal Teknologi

CRUDE OIL BIOREMEDIATION BY INDIGENOUS BACTERIA ISOLATED FROM OILY SLUDGE

Nur Hafizah Azizan^o, Kasing Ak Apun^b, Lesley Maurice Bilung^b, Micky Vincent^b, Hairul Azman Roslan^b, Awang Ahmad Sallehin Awang Ahmad Husaini®

Department of Biotechnology, Kulliyyah of Science, International Islamic University Malaysia, Kuantan Campus, Jalan Istana, Bandar Indera Mahkota, 25200, Kuantan, Malaysia Department of Molecular Biology, Faculty of Resource Science

and Technology, University Malaysia Sarawak, 94300, Kota Samarahan, Sarawak, Malaysia

Graphical abstract Abstract



ulture technique leads to the discovery of six presumptive TPH-ntfliction and characterization tests using mapphological, bioche chniques have successfully isolated Pseudomonas aeruainas accers (UMAS2SF) and Nebbiella spp. (UMAS3RF). All strains were o Mattin managements (unreaser) and energy source for their unde oil as sole carbon and energy source for their Minimal Salt medium supplemented with 1% (v/v) rey produced the highest cell counts on the third

NUSANTARA BIOSCIENCE Vol. 13, No. 1, pp. 1-10 May 2021

Potential of Candida glabrata from ragi as a bioethanol producer using selected carbohydrate substrates

MICKY VINCENT, OUEENTETY JOHNNY, DAYANG SALWANI AWANG ADENI, NURASHIKIN SUHAILI cience and Technology, Universiti Malaysia Sarawak. 94300 Kota Samarah Tel.: +60-825-82985, Fax.: +60-825-83160, •email: vmicky@unimas.my

Manuscript received: 16 September 2020. Revision accepted: 1 December 2020.

biorefinery context

Potential of sugar beet vinasse as a feedstock for biocatalyst production within an integrated

Nurashikin Suhaili,^{a,b} Max Cárdenas-Fernández,^a John M Ward^a and Gary J Lye^{a*}

Abstract

BACKGROUND: This work explores the feasibility of vinasse as an inexpensive feedstock for industrial biocatalyst production within the context of an integrated sugar beet biorefinery. As an exemplar, production of CV2025 @-Transaminase (@-TAM) in Estencifica coll BL21 was studied.

In Escherichia coli BL21 was studied. RESULTS: Characterisation of vinasse showed that it comprised mainly of glycerol along with several reducing sugars, sugar alcohols, acctate, polyphenolis and protein. Preliminary results showed E. Coli BL21 cell growth and CV2025 m⁻²Am production were feasible in cultures using 17% to 25% (v/v) vinasse with higher concentrations demonstrating inhibitory effects. The orgalactose present in vinasse facilitated auto-induction of the pQR801 plasmid enabling (ZV2025 m⁻²Am production addition of expensive isopropyl-*β*-o-thiogalactopyranoside (IPTG). Assessment of different vinasse pre-processing options confirmed simple dilution of the vinasse was sufficient to reduce the concentration of polyphenols to below inhibitory levels. Optimisation experiments, carried out using a controlled, 24-well microbioreactor platform, showed supplementation of diluted vinasse medium with 0 gL⁻¹ speat extract enabled enhancements 0 z.8, 25, 5, 4 and 3-fold in specific growth rate, maximum biomass concentration, CV2025 m⁻²Am volumetric and specific activity, respectively. Investigation into the metabolic preferences of E. Coli BL21 when grown in vinasse showed a preference for D-mannitol utilisation before simultaneous solver a 200-fold volumetric case leat translation and maxtate dassed on matched volumetric mass transfer coefficient (k, a). The results showed good comparability with respect to cell growth, substrate consumption and CV2025 m⁻²Am production representing operated at higher k, a values. CONCLUSION: This work describes the promision andiration of ulaxes for another the simulation before the simulation set of the simulation before the simulation presenting approximation of plate translation and active and substrate consumption and CV2025 m⁻²Am production representing operated at higher k, a values. CONCLUSION: This work describes the promision andiration of ulaxes for another data data before the simulation before the simulation before the simulation befo

CONCLUSION: This work describes the promising application of vinasse for production of microbial enzymes and insights into carbon source utilisation in complex feedstocks. Exploitation of vinasse as a fermentation feedstock could be further extended to other processes involving different microorganisms and target enzymes. © 2018 The Authors. Journal of Chemical Technology & Biotechnology published by John Wiley & Sons Ltd on behalf of Society of Chemical Industry.

Supporting information may be found in the online version of this article

Keywords: E. coli BL21 metabolism; integrated biorefinery; microbioreactor; sugar beet vinasse; transaminase

INTRODUCTION

INTRODUCTION Winsses is the stillage released following distillation of yeast fer-mentation broth during bioethanol production form either sugar-cane or sugar beet.¹ It represents one of the main waste streams discharged by bioerfenreis in India, South America and Europe, which normally integrate sugar manufacturing plants with ethanol distilleries. In Brazilian and Indian bioerfenreis for example, the average volumetric production of vinasse is reported to be 10 to 15 fold greater than the ethanol generated.³³ Currently, vinasse is mainly exploited for soil mineralisation and as an additive for fertilisers and animal feeds due to the high content of organic nutrients.⁴ However, the market for these applications is not predicted to grow at the same rate as for biofuel production. A large amount of vinasse is still

disposed of into water streams, from some manufacturing sites causing an adverse impact on the aquatic ecosystem due to the presence of toxic compounds.⁵ The increasing production of vinasse, as a consequence of the growing demand for bioethano

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a Department of Biochemical Engineering, The Advanced Centre for Biochemical Engineering, University College Landon, London, UK b Faculty of Resource Science and Technology, UNIMAS, Kota Samarahan,

earch Journal of Biotechnology

Vol. 17 (7) July (2022) Res. J. Biotech

Autolysis of spent baker's yeast generated from sago bioethanol fermentation: a preliminary study

Mahmod Nik Nur Aziati, Suhaili Nurashikin*, Awang Husaini Awang Ahmad Sallehir and Awang Adeni Dayang Salwani Faculty of Resource Science and Technology. UNIMAS, 94300 Kota Samarahan, Sarawak, MALAYSIA "unrashikini?mainas.uw"

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> nt Baker's yeast (Saccharomyces cerevisiae) is tined as a by-product of ethanol fermentation istry and is mainly discarded as waste. The spent st can be potentially converted to yeast extract rugh autolysis process in which the yeast cells are d under certain conditions to release valuable and active substances. To date, there are still limited terve substances. To date, there are stull limited lies on the autolysis of spent S. cerevisiae generated wing the production of sago bioethanol. This arch aims to investigate the feasibility of autolysis pent Baker's yeast generated from sago bioethanol sentation by focusing on two important parameters sely initial pH and incubation time. The spent er's yeast was autolysed at different initial pH ues (3, 5 a

AsPac J. Mol. Biol. Biotechnol. 2020 Vol. 28 (3) : 44-51

Keywords: Autolysis, Bioethanol fermentation, Saccharomyces cerevisiae, Spent Baker's yeast, Yeast Keywords: extract.

Introduction

Baker's years (Saccharomyces cerevisiae) is commonly utilised in the fermentation industry and the production generates Large quantities of spent yeast as their by-products^{9,24}. About 400 000 tons of spent Baker's yeast are produced as a by-product by the brewing industry annually. The waste, in general, is still underused and is normally discarded into the environment⁷. The spent yeast is a rich source of proteins, essential amino acids, RNA, vitamins B and minerals which essentially serve as a raw material in the production of yeast extract³³. The extracts from the spent yeast can serve as promising sources of valuable ingredients

Enhancement of very high gravity bioethanol production via fed-batch fermentation using sago hampas as a substrate

Nur Adila Muradi, Dayang Salwani Awang Adeni", Nurashikin Suhaili

ere Biotechnology Programme, Faculty of Resource Science and Technology, Universiti Malaysia Sarawak, 94300 Kota Samaraban, Sarawak, Malaysia

Received 27th February 2020 / Accepted 10th June 2020

Abstract. Very high gravity (VHG) ethanolic fermentation is a promising technology used for producing bioethanol. However, the technology is often associated with the excessive amount of glucose that is entirely supplied in the beginning of the culture causing the fermentation process to be sluggish and Abstract. Vincent M, Johnny Q, Adeni DSA, Subaili N. 2021. Potential of Candida glabrata from ragi as a bioethanol producer using the fermentation process to be sluggish and selected carbohydrate substrates. Nusantara Bioscience 13: 1-10. The flexibility and efficiency of fermenting microorganisms to convert medium also elevates the comotic pressure, which has a destructive effect on yeast cells. This study arms to enhance the production of VHG bioethanol from sago hampas hydrolysate (SHH) via fed-batch fermentation was or enhance the production of VHG bioethanol from sago hampas hydrolysate (SHH) via fed-batch from the object and control. Our results showed that the maximum yeast cell concentration achieved was from ragi as a bioethanol producer using the fermentation. The fermentations were performed in a 2-1 stirred tank bioreactor. Batch fermentation was of from sago hampas hydrolysate (SHH) via fed-batch sage and complex of the object via from sage as a control. Our results showed that the maximum yeast cell concentration achieved was vield atta

Malaysian Journal Of Science 40(3): 43-53 (October 2021)

THE CHARACTERISTICS OF SAGO FROND SAP FROM TWO SELECTED GROWTH STAGES; ANGKAT PUNGGUNG AND UPONG MUDA PALMS

Nurazureen Matnin^{1a}, Dayang Salwani Awang Adeni^{2a*}, Muhammad Norhelmi Ahmad^{3a} and Nurashikin Suhaili^{4a}

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*Corresponding Author: adsalwa@unimas.my Received: 13th Apr 2021 Au DOI: https://doi.org/10.22452/mjs.vol40no3.4 Accepted: 2nd Aug 2021 Published: 31st Oct 2021

ABSTRACT Sago frond is produced in abundance upon harvesting of the sago palms for starch extraction, hence need to be utilized and developed into beneficial products. In this study, the sap which contains sugars and starch is obtained by roller crushing the skinned frond for use as fermentation medium. Fronds from two selected growth stages (namely Angkat punggung and Upong muda) and two different positions within the rosette (inner and outer circle) of the sago palm were studied. Based on the results, the outer circle frond of Upong muda palm gave the highest volume of sap at 290mL/kg which equivalent to 1600 mL/frond. On top of that, sago frond sap has an acidic pH, with glucose as major sugar component and contained various kinds of minerals like calcium, potassium and manganese. All fronds from two selected growth stages contain glucose between 28-68 g/L and xylose 21-29 g/L, respectively. After 21 days of storage, it can be concluded that the amount of reducing sugars and starch in most samples obtained from two selected growth stages decreased slightly from the original. Subsequently after this study, both fresh and stored sago frond sap can be used as a fermentation substrate without any pre-treatment or modification.

Keywords: Angkat punggung, fermentation, starch, sugar, Upong muda.

MEET OUR EXPERTS NICHE: GENETICS



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Biotechnology has revolutionized genetics study by providing powerful tools for genome analysis, genetic engineering, functional genomics, personalized medicine, evolutionary genetics, and conservation biology. These advancements have deepened our understanding of genes, their functions, and their impact on health, disease, and biodiversity, leading to breakthroughs in various scientific fields and potential applications in healthcare, agriculture, and environmental conservation.

NICHE: GENETICS

SOME OF OUR RECENT WORKS



Lytogenetic, chromosome count optimization and automation of Neolamarckia cadamba (Rubiaceae) root tips derived from *in vitro* mutagenesis

Wee-Hiang ENG¹, Wei-Seng HO^{1*}, Kwong-Hung LING²

¹Universiti Malaysia Sarawak, Forest Genomics and Informatics Laboratory (IGiLab), Faculty of Resource Science and Technology 94300 Kota Samarahan, Sarawak, Malaysia; engweehiang@gmail.com; wsho@unimas.my ('corresponding author) ²Sarawak Timber Association, Wisma STA, 26, Jalan Datuk Abang Abdul Rahim, 93450 Kuching, Sarawak, Malaysia; khling68@gmail.com

Abstract

Chromosome count is the only direct way to determine the number o study is often considered trivial that seldom described and discussed in deta the chromosome count protocol should be revised and revisited before it bec study, we encountered challenges in obtaining a clear micrograph for the chro cells of *Neolamarckia cadamba* (Roxb.) Bosser (Rubiaceae) root tips. Set through micrograph observation, such as existing unwanted particles in cells chromosome clumping. To overcome these, root tip types, staining method



Chiang Mai J. Sci. 2017; 44(4) : 1304-1310 http://epg.science.cmu.ac.th/ejournal/ Contributed Paper

Sequence Characterized Amplified Region Markers for Species-specific Identification of Three Threatened Aquilaria Species

Hairul Azman Roslan*[a], Md. Anowar Hossain [a,c], Nur Qistina Othman [a,b], Cheksum Supiah Tawan [b] and Isa Ipor [b]

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> Received: 26 January 2016 Accepted: 13 June 2016

Current Applied Science and Technology Vol. 20 No. 2 (May-August 2020)

Sago Palm Genome Size Estimation via Real-Time Quantitative PCR

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Received: 9 December 2019, Revised: 23 February 2020, Accepted: 27 February 2020

Abstract

Sago palm, *Metroxylon sagu* Rottb., is an underutilized indigenous food crop that can be found mainly in the South East Asia and Pacific regions. It is a main starch producer and socioeconomically important crop in the South East Asia region including Malaysia. The sago starch provides for considerable potential to food security in the places where it is grown. However, not many molecular works have been reported thus far. In the post genomic era, sago plant genome sequencing is very important for sustainable starch development in these regions. Therefore, determination of the genome size is prerequisite to full genome sequencing and assembly. Here we report on the use of real-time quantitative polymerase chain reaction (qPCR) in determining the genome size. For this work, we calculated the genome size, (bp) of *M. sagu* based on qPCR-derived copy number of two single copy genes. *Pichia pastoris*, with a known genome size of *M. sagu* was calculated to be 1.87 Gbp. This genome size information would be beneficial for subsequent molecular work with including genome sequencing and analysis on this economically important crop plant.

Keywords: Genome size, Metroxylon sagu, real-time PCR, copy number, Pichia pastoris DOI 10.14456/cast.2020.10



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NICHE: MOLECULAR BIOLOGY

SOME OF OUR RECENT WORKS

		Research	Article	
Food Constant	APPLIED FOOD BIOTECHNOLOGY, 2020, 7 (3):127-134 Journal homepage: www.journals.sbmu.ac.ir/afb	pISSN: 234 efSSN: 241	15-5357 23-4214	
Toxicity Asses	ssment of Lactococcus lactis IO-1 Used in Co	conut Bevera	ages	
against Artem	ia salina using Brine Shrimp Lethality Test		Der Springer Link	
Dlaide Olawunmi A Wahab Abideen‡, Fo	jibola1°, Samuel Lihan², Ahmad Hussaini1, Rosmawati Saat1, azia Mohamad Sinang1, Ngieng Ngui sing1, Gbadebo Clement	ldris Adewale Ahı Adeyinka ^s	Home > Advances in Traditional Medicine > Article	
- Faculty of Resource Sci	ience and Technology, Universiti Malaysia Sarawak, Kota Samarahan, 94300, Ma	laysia.		
- Institute of Biodiversity	v and Environmental Conservation, Universiti Malaysia Sarawak, Kota Samarahan	, 94300, Malaysia.	Short Communication Published: 30 September 2020	
- Center for Natural Prod	Justs Research and Drug Discovery, University of Malaysia, 50603, Kuala Lumpu	, Malaysia.	Anticancer and antimicrobial peptides from medicinal	
- Department of Stateook	ongy, Facury of nanc and Appued Science, Orun State University, Osogoo, Ng al Engineering, Mangosuthu University of Technology, 51, Umlazi, Durban, South	Africa.	plants of Borneo island in Sarawak	
Abstract		Article Informa	Hasnain Hussain ⊠, Nurhazlina Hamdan & Edmund Ui-Hang Sim	
Background and o	biective: Plant-based fermented foods containing favorable micro-	Article history:	Advances in Traditional Medicine 21, 189–197 (2021) Cite this article	
organisms have been compounds that are I appropriate assay to	in used to improve diets. Starter microorganisms may produce toxic hazardous to consumers. Brine shrimp lethality test is a convenient and check toxicity of camdos. The aim of this study was to investigate	Received 09 M Revised 12 Ap Accepted 20 Ap	290 Accesses 4 Citations Metrics	
Current App	plied Science and Technology Vol. 20 No. 3 (September - December 20)20)	Abstract	
		- 1	The interest in drug discovery from plants-based metabolites has been of interests to	
Ennueston	of Decembinant Alashal Dahudusan assin Fach	michia	researchers, especially for health well-being and for therapeutic reasons. The work described	
Expression	of Recombinant Alconol Denyarogenase in Esch	ericnia	here was to explore the in vitro anticancer and antimicrobial peptides from six traditional	
con s	Transformation of Tomato Soads		medicinal plants commonly used in Sarawak, Malaysian Borneo. Proteins were extracted from	
	Transformation of Tomato Seeus		plants with a common protein extraction buffer. Evaluation for in vitro anticancer activity was	
	Mastura Sani ^{1*} and Hairul Azman Roslan ²		done against normal and carcinomas nasopharyngeal cell line; NP69 and HK-1 respectively by	
			using established MTT microtiter plate assays. Antimicrobial activity was tested against	
Food Technology, School of Engineering and Technology, University College of		Staphylococcus aureus and Escherichia coli, a Gram-positive and Gram-negative respectively.		
² Ecoulty of Pos	Technology Sarawak, Sarawak, Malaysia	arouvals		
Faculty of Ke	Malaysia	arawak,		
Received	1: 26 December 2019, Revised: 13 March 2020, Accepted: 24 April 202	20	Penyelidik UKM dan UNIMAS Hasilkan Sistem	
			Penskoran Madu Kelulut UKSF	
	Abstract		13/09/2022 / Admin	
Alcohol dehydrog	genase is an enzyme that is involved in various roles in plant such	h as plant	al Post Views: 638	
development, grov dehvdrogenase 1 (wth and plant responses to abiotic and biotic stresses. A recombina (Adh1) cDNA (r-msAdh1) from Metroxylon sagu has been previously isol	nt <i>alcohol</i> ated, and it		
contained 20 nucle	eotides derived from Elaeis guineensis at the 5'-end and had a molecular	weight of		
1.14 kb. The obj	ective of this study is to determine the function of r-msAdh1 via a	nalyses in		
		SN 2586-9000		
AH	P-IS: E-IS:	SN 2586-9027		
	Homepage : https://tci-thaijo.org/index.php/SciTechAsia Science & Tec	hnology Asia		
47	Vol.26 No.2 April - June 2021 Pr	age: [155-168]		
	Original research article			

Effects of Short Term UVB and UVC Irradiation on Hydroxyphenylpyruvate **Reductase Expression and Rosmarinic Acid** Accumulation in Orthosiphon aristatus

Hairul Azman Roslan^{1,*}, Felicia Chin Yien Yin¹, Sim Siong Fong¹, Alan Fong², Johnson Chong3, Ahmad Husaini1

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ABSTRACT

Ultraviolet (UV) light is one of the abiotic stresses experienced by plants. Generally, UV irradiation can affect the growth and development of plants and radiation above ambient level can cause the activation of plant stress responses through the self-protective secondary metabolism system. One of the products of these responses is rosmarinic acid, which is an important secondary metabolite in the medicinal plant, Orthosiphon aristatus. Rosmarinic acid exhibits many pharmacological and biological properties such as antioxidant, anticancer, antiviral, antibacterial, and anti-inflammatory activity, as well as other health promoting



Oleh Siti Aishah Mohd Yusot Foto Ikhwan Hashim

BANGI 12 September 2022 - Sekumpulan penyelidik dari Universiti Kebangsaan Malaysia (UKM) dan Universiti Malaysia Sarawak (UNIMAS) berjaya menghasilkan satu inovasi hakcipta sistem penskoran bagi produk madu kelulut iaitu Unique Kelulut SuperFood Factor (UKSF).

Sistem penskoran madu kelulut UKSF itu dapat membantu meningkatkan lagi kualiti madu kelulut Malaysia ke peringkat antarabangsa.

Timbalan Naib Canselor Hal Ehwal Penyelidikan dan Inovasi, Prof. Dato' Dr. Wan Kamal Mujani berkata sistem penskoran yang dihasilkan itu juga boleh diguna pakai bagi semua produk madu kelulut yang dihasilkan kerana tidak terhad kepada Malaysia sahaja, malah kepada Indonesia, Brazil dan beberapa negara pengeluar lain di dunia

MEET OUR EXPERTS NICHE: ENZYMOLOGY



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Enzymes are widely utilized in biotechnological processes and industries. They serve as catalysts for chemical reactions, enabling the production of various valuable products such as biofuels, pharmaceuticals, and biodegradable plastics. Enzymes also facilitate efficient and environmentally friendly industrial processes by reducing energy consumption and waste generation.

NICHE: ENZYMOLOGY

SOME OF OUR RECENT WORKS

D Springer Link

Home > Brazilian Journal of Microbiology > Article

Biotechnology and Industrial Microbiology - Research Paper | Published: 16 September 2022

Characterization of bioactive compounds produced by endophytic fungi isolated from *Gynura procumbens* (Sambung Nyawa)

Haifa Arghnia A. Jamal, Ahmad Husaini 🗁, Ngieng Ngui Sing, Hairul Azman Roslan, Azham Zulkharnain & Wahab Abideen Akinkunmi

Brazilian Journal of Microbiology 53, 1857–1870 (2022) Cite this article

196 Accesses | 1 Citations | 2 Altmetric | Metrics

- A <u>Correction</u> to this article was published on 15 November 2022
- 3 This article has been updated

Abstract

This research evaluates the bioactivity of twelve endophytic fungi successfully isolated and characterised from *Gynura procumbens*. The fungal extracts displayed inhibitory activity against *Staphylococcus aureus*, *Pseudomonas aeruginosa*, Methicillin-resistant *Staphylococcus aureus* (MRSA), *Escherichia coli* and *Salmonella typhi* with the MIC and MBC of 5000 µg/mL. High antioxidant activity using DPPH free radical scavenging assay with inhibition of 86.6% and IC_{50} value of 104.25 ± 18.51 µg/mL were exhibited by ethyl acetate



Protein Expression and Purification Volume 164, December 2019, 105462



Characterization and expression in *Pichia pastoris* of a raw starch degrading glucoamylase (GA2) derived from *Aspergillus flavus* NSH9

Kazi Muhammad Rezaul Karim^a, Ahmad Husaini^b A, 🔯 , Ngieng Ngui Sing.^b, Tasmia Tasnim^c, Fazia Mohd Sinang.^b, Hasnain Hussain^b, Md Anowar Hossain^d, Hairul Roslan^b

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dilus fisme NCUO sees one

https://doi.org/10.1016/j.pep.2019.105462 >

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Abstract

Research Article

Decolourisation Capabilities of Ligninolytic Enzymes Produced by *Marasmius cladophyllus* UMAS MS8 on Remazol Brilliant Blue R and Other Azo Dyes

ng a pH and thermostable glucoamylase with a ed in <u>Pichia</u> pastoris to produce recombinant oamylase gene (2039 bp), and cDNA (1839 bp) nilar to glucoamylase from <u>Aspergillus oryzae</u> ned to be a <u>signal peptide</u> for secretion, and the

Ngieng Ngui Sing, Ahmad Husaini, Azham Zulkharnain, and Hairul Azman Roslan

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Received 27 July 2016; Accepted 20 December 2016; Put to the second

Academic Editor: Dong-sheng Shen

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Marasmius cladophyllus was examined for its ability t (RBBR) and screened for the production of ligninolyti decrease in absorbance ratio of A_{592}/A_{500} shows that Marasmius cladophyllus produces laccase and lignin per activities were increased, with laccase activity recorded dye decolourisation using the culture medium shows th 3 Biotech (2018) 8:204 https://doi.org/10.1007/s13205-018-1225-z

ORIGINAL ARTICLE

CrossMark

Purification of an alpha amylase from Aspergillus flavus NSH9 and molecular characterization of its nucleotide gene sequence

Kazi Muhammad Rezaul Karim^{1,2} - Ahmad Husaini¹ · Ngieng Ngui Sing¹ - Fazia Mohd Sinang¹ · Hairul Azman Roslan¹ - Hasnain Hussain¹

Received: 23 November 2017 / Accepted: 22 March 2018 / Published online: 29 March 2018 © Springer-Verlag GmbH Germany, part of Springer Nature 2018

Abstract

In this study, an alpha-amylase enzyme from a locally isolated *Aspergillus flavus* NSH9 was purified and characterized. The extracellular α -amylase was parified by ammonium sulfate precipitation and anion-exchange chromatography at a final yield of 2.55-fold and recovery of 11.73%. The molecular mass of the purified α -amylase was estimated to be 54 kDa using SDS-PAGE and the enzyme exhibited optimal catalytic activity at pH 5.0 and temperature of 50 °C. The enzyme was also thermally stable at 50 °C, with 87% residual activity after 60 min. As a metalloenzymes containing calcium, the purified α -amylase showed significantly increased enzyme activity in the presence of Ca²⁺ ions. Further gene isolation and characterization shows that the α -amylase gene of *A. flavus* NSH9 contained eight introns and an open reading frame that encodes for 499 amino acids with the first 21 amino acids presumed to be a signal peptide. Analysis of the deduced peptide sequence showed the presence of three conserved catalytic residues of α -amylase, two Ca²⁺-binding sites, seven conserved peptide sequences, and several other properties that indicates the protein belongs to glycosyl hydrolase family 13 capable of acting on α -1,4-bonds only. Based on sequence similarity, the deduced peptide sequence of *A. flavus* NSH9 α -amylase was also found to carry two potential surface/secondary-binding site (SBS) residues (Trp 237 and Tyr 409) that might be playing crucial roles in both the enzyme activity and also the binding of starch eranules.



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