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O-1

Tissue-based proteomics: insight into molecular mechanisms in cervical carcinogenesis

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Background

Tissue-based proteomics is an evolving tool used in cancer research to characterize the pathophysiology of disease. However, the proteome alterations involved in cervical carcinogenesis are not extensively studied. This study aims to elucidate the differentially expressed proteins and offer insights into the cellular processes and pathways involved in the development of cervical cancer.

Methodology

The pathological regions of interest in the cervical squamous epithelium were micro-dissected from formalin-fixed paraffinembedded (FFPE) tissue sections of six normal cervix cases, five HPV-associated squamous intraepithelial lesion (SIL), and six squamous cell carcinomas (SCC). The samples were trypsin digestion and subjected to high throughput liquid chromatography-

electrospray ionization-tandem mass spectrometry (LC/ESI-MS/MS) and trapped ion mobility time-of-flight-mass spectrometry (tim-sTOF-MS), followed by quantification with MaxQuant and Perseus software. Bioinformatics analyses were carried out using DAVID, ConsensusPathDB, and STRING.

Results and Discussion

We identified a total of 3597 proteins with 589, 550, and 1570 proteins unique to the normal cervix, SIL, and SCC groups, respectively, while 332 proteins were similar across all three groups. The predominant protein found was histone. Interestingly, the quantification results showed an upward trend for the up-regulated proteins and a downward trend for the down-regulated proteins in the progression from normal to SIL and SCC. The main molecular function was the binding process, and the top biological processes were chromatin silencing for SIL compared to the normal cervix and nucleosome assembly for the SCC compared to SIL group. The key pathways involved were viral carcinogenesis and necroptosis, reflecting their role in cell proliferation, migration, and metastasis.

Conclusion

The identification of proteins and their associated pathways provides a deeper understanding of the underlying molecular mechanisms involved in HPV-associated cervical cancer.

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Page 2 of 46

0-2

The expression of apoB and 4HNE in overweight-and obeserelated colorectal carcinoma tissues

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Background

Obesity has been found to be related to an increase in the incidence and progression of colorectal carcinoma (CRC) [1]. However, the relationship between obesity and CRC is still unclear. Low-density lipoprotein (LDL) is known for causing abnormal lipid metabolism and is usually measured in the blood [2]. Two LDL-related biomarkers, apoB and 4HNE, were chosen to investigate their expression in overweight-and obese-related CRC tissues, and their association with the clinicopathological data was also determined.

Methodology

Human ethical approval with a series code of USM/JEPeM 19060354 was obtained. This retrospective study involved overweight-and obese-related CRCs diagnosed in HUSM from January 2015 to December 2021. The classification of BMI followed the Asia Pacific BMI classification [3]. Clinicopathological data was retrieved from Laboratory Information System (LIS) and medical records at Pathology laboratory, HUSM. The CRC archival tissue blocks were then collected for Immunohistochemistry (IHC) staining. The data was then analysed using SPSS Version 26.

Results and Discussion

A total of 69 overweight-and obese-related CRCs were retrieved within seven years. The patients had median age of 61 years old with interguartile range of 21, common in age of more than 50 years old (73.9%) and in males (52.2%). Most of the patients were found with CRCs localised in sigmoid colon (30.4%) and rectosigmoid colon (24.6%), larger than 2cm of tumour (97.1%), presence of bowel wall invasion (94.2%), and absence of lymph nodes involvement (58.0%). The tumours were frequently classified as adenocarcinoma (91.3%) with moderate differentiation grade (82.6%) and under Modified Duke B class (55.1%). The IHC-stained CRC tissue slides showed high apoB expression (91.3%), whereas low 4HNE expression (82.6%). Significant associations were observed between apoB expression with age (p value=0.036), tumour site (p value=0.048), bowel wall invasion (p value=0.035), Duke classification (p value=0.002), while 4HNE expression with tumour size (p value=0.029). Majority of CRC tumours of more than 2cm showed low 4HNE expression. This may be explained by the protective effect of low 4HNE level that prevents the damage of the cancer cells [4].

Conclusion

In conclusion, the contrast tissue expression of apoB and 4HNE, and the significant associations obtained has shed light on the role of these LDL-related biomarkers in overweight-and obese-related CRC tissues. Exploration on the role of these biomarkers is recommended for *in vitro* study.

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O-3

Quantitative proteomics analysis of insecticide resistant *Ae. aegypti* Abubakar Shettima^{1, 2}, Intan H Ishak^{3, 4}, Benjamin Lau⁵, Hadura Abu Hasan^{3,4} and Noorizan Miswan⁶ and Nurulhasanah Othman¹
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Background

Synthetic insecticides are the main vector control method globally. However, the widespread use of insecticide is causing resistance in mosquitoes. Understanding the proteomics basis of insecticide resistance may provide novel opportunities to control mosquito vectors. Hence, this study aimed to elucidate proteins associated with permethrin resistant *Ae. aegypti* using quantitative proteomics.

Methodology

The study evaluated the resistance pattern of *Ae. aegypti* from dengue hotspot and non-hotspot areas of Penang Island. Permethrin 0.75% insecticide-impregnated papers were used to determine the resistance status in adult female *Ae. aegypti*. The mosquito proteins were analysed using LC–ESI–MS/MS for protein identification and quantification via label-free quantitative (LFQ) analysis. In this study, differential protein expression (DEP) analysis was carried out using Perseus 1.6.14.0 statistical software to perform ANOVA and student's t-test. Protein-protein interaction (PPI) and functional ontology enrichment analyses were performed using STRING software.

Results and Discussion

The bioassay results showed 28% and 53% mortalities in mosquitoes exposed to permethrin from the hotspot and non-hotspot areas. Permethrin has been used extensively in Malaysia's dengue vector control program. The mortality percentage has shown a high permethrin resistance rate in the field strain Ae. aegypti mosquito, including the strains from the dengue hotspot and non-hotspot areas. These results suggested high resistance status in the dengue hotspot and non-hotspot areas. The high resistant patterns identified in the field strain Ae. aegypti could be because of the widespread and indiscriminate permethrin use in the investigated areas, especially if a site was labeled as a dengue hotspot. The resistance in the non-hotspot area could be because the area was once a dengue hotspot. These areas received extra attention from vector control management and private contractors lead to insecticide selection pressure and produce offspring carrying insecticide-resistant genes. The LFQ analyses revealed 501 (q-value <0.05) DEPs. The t-test showed 114 upregulated and 74 downregulated proteins in resistant versus laboratory strain exposed to permethrin. The significant functional ontology enrichment of the DEPs indicated drug-metabolic process, small molecule metabolic process, hydrolase activity acting on ester bonds, and catalytic activity. The PPI of DEPs showed a p-value at <1.0 x 10⁻¹⁶ in permethrin-resistant *Ae. aegypti.* Significantly enriched pathways in DEPs revealed metabolic pathways, oxidative phosphorylation, carbon metabolism, biosynthesis of amino acids, glycolysis, and citrate cycle.

Conclusion

This study has revealed several DEPs and highlighted upregulated and downregulated proteins associated with insecticide resistance in Ae. aegypti.

0-4

Clarithromycin resistance in *Helicobacter Pylori* is associated with genetic polymorphism of virulence factors

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Background

For a long time, the macrolide antibiotic clarithromycin was used to treat *Helicobacter pylori* (*H. pylori*) infections. The efficiency of traditional triple therapy for the eradication of *H. pylori* has recently been dismally reduced in many nations [1]. Clarithromycin resistance is spreading over the world and is the leading cause of *H. pylori* treatment failure [2]. We employed whole-genome sequencing (WGS) technology to identify genomic alterations related with the development of antibiotic resistance in *H. pylori* with induced resistance to Clarithromycin.

Methodology

The clarithromycin-sensitive *H. pylori* strains were induced to become resistant against clarithromycin. To induce resistance, the strains were exposed to gradually increased concentration of clarithromycin in vitro for 3-5 days on a chocolate agar plate (CAP). The identity between resistant strains and their corresponding parental sensitive strains before induction were verified by random amplification of polymorphic DNA polymerase chain reaction (RAPD-PCR). WGS for bacterial DNA was performed, using Illumina platform.

Results and Discussion

There were 113 distinct single nucleotide variants (SNVs) and 286 unique insertions and deletions (InDels) changes discovered among the induced-resistant *H. pylori* strains. Interestingly, among the induced-resistant *H. pylori* strains, mutations in *cag1*, *cag4*, *fliR*, *obgE*, *tlpA*, and *vacA* were detected, which are known virulence genes that may help in bacterial survival, indicating that those mutations may be associated with the emergence of resistant *H. pylori* [3]. Moreover, we also found that it is not possible to infer a Clarithromycin resistance phenotype based on the existence of distinct point mutations in A2143G and/or A2142G mutations in 23S rRNA and may not be sufficient for predicting Clarithromycin resistance in *H. pylori*.

Conclusion

As a result, we hypothesise that in these strains, alternative mechanisms unrelated to the 23S rRNA gene sequence, such as the existence of an efflux pump, may play a role in Clarithromycin resistance.

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0-5

Metabolome analysis of induced-resistance *Helicobacter pylori* against Clarithromycin

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Background

Helicobacter pylori (H. pylori) is a gram-negative bacterium that thrives in stomach mucus and epithelial mucosa, causing gastric ulcers which may develop into gastric cancer. One of the most prevalent causes of treatment failure is the emergence of antibiotic-resistant of H. pylori infection [1]. The goal of this work is to use a metabolomic method to discover the metabolites associated with Clarithromycin-resistance in H. pylori.

Methodology

Clarithromycin-sensitive *H. pylori* strains were induced to become resistant against Clarithromycin. To induce resistance, the strains were treated to 0.1x, 0.25x, 0.5x, 1X, 2x, 4x, 8x, 16x, and 32x MICs of Clarithromycin in vitro for 3-5 days. Bacterial metabolites were isolated using the Bligh and Dyer technique and analysed using liquid chromatography-mass spectrometry (LCMS) [2]. The MassHunter Qualitative Analysis and Mass Profiler Professional tools were used to process and analyse the data. Sensitive, pre-resistant, and induced-resistant are three separate categories based on global metabolomic patterns

Results and Discussion

Using one-way ANOVA, a total of 982 molecular features were identified to be significantly different (p-value < 0.005) between sensitive, pre-resistant, and resistant H. pylori strains. Additionally, 432 molecular features matched the metabolites in the Agilent METLIN Accurate Mass-Personal Metabolite Database and Library (AM-PCDL) database based on accurate mass, isotope ratios, abundances and spacing. In contrast to sensitive strains, induced-resistant strains generated more metabolites (585 features). Further investigation was carried out in order to find metabolites that varied substantially (p-value <0.05) between sensitive, pre-resistant, and induced-resistant H. pylori strains. These metabolites include lipids and metabolites involved in the metabolism of fructose and mannose. Our data imply that D-Mannitol, L-Leucine, and Pyridoxine, which correlate with bacterial survival, may constitute a potential antibiotic mechanism [3].

Conclusion

Understanding the underlying metabolic variations between Clarithromycin-sensitive *H. pylori* strains and Clarithromycin-resistance *H. pylori* strains may be a promising technique for developing novel antibiotic candidates.

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Page 4 of 46

0-6

The antimicrobial effect of vinegar on Escherichia coli O157:H7 isolated from lettuce

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Background

Lettuce (Lactuca sativa) is a ready-to-eat (RTE) vegetable, which is popular among consumers due to its convenience and high nutritional value. Unfortunately, lettuce is frequently associated with Escherichia coli O157:H7 outbreaks that cause serious illnesses such as haemolytic uremic syndrome and haemorrhagic colitis [1]. Thus, a proper cleaning of the products is crucial to eliminate the risk of foodborne pathogens contamination. Acetic acid, such as vinegar, has shown its antimicrobial effect on inhibiting the growth of E. coli O157:H7 [2]. In addition, it is commonly used in the household settings for cooking and washing. Hence, the objectives for this study were identifying and verifying the presence of E. coli O157:H7 on lettuces purchased from retail market in Kampar Perak, using multiplex polymerase chain reaction (mPCR); followed by evaluating the antimicrobial activity of vinegar based on the analysis of log reduction in growth of E. coli O157:H7.

Methodology

A total of 20 samples of lettuce was purchased from a retail market in Kampar Perak. Homogenization was performed on 10 g of shredded lettuce, followed by incubation at 37°C for 24 h in Tryptic Soy Broth (TSB) to enrich the bacteria. Serial dilution was performed prior to plating on selective agar for *E. coli* O157:H7. Verification of colonies was done using multiplex polymerase chain reaction (PCR) with primer pairs targeting genes of rfb_{O157} (256 bp) and $flic_{H7}$, (625 bp). Both encodes for somatic and flagellar antigen, respectively. On the other hand, 1.0 McFarland ATCC 700728 *E. coli* O157:H7 was prepared and inoculated on lettuce at 5°C for 1 h. 5% of white vinegar was immersed into the 'contaminated' lettuce at 4 time points:10, 20, 30 and 40 mins. Antibacterial effect was evaluated by enumerating the reduction of bacteria colonies in log CFU/g. The significance of log reduction of inoculated bacteria was analysed quantitatively using One-Way ANOVA test (Graph Pad Prism 9.0) with 95 % confidence level.

Results & Discussion

None of the samples collected were found contaminated with *E. coli* O157:H7. This might be due to the safety measures taken by the Food Safety and Quality Division (FoSIM) in maintaining the quality of products in the retail market. However, in this study, 5 samples of lettuce (25%) showed presence of $flic_{H7}$ gene but not rfb_{O157} virulence gene. This might postulate the presence of other *E. coli* serotypes such as O55:H7 which shares similarity in flic H7 gene. On the other hand, significant reduction of *E. coli* O157:H7 was observed after 40 min treatment with 5% vinegar. Organic acids exert their antimicrobial effect by altering the pH level of bacteria's cytoplasm which then leads to cell lysis due to the disruption of enzymes' and proteins' structures [3]. Hence, acetic acid is the best alternative to chlorinated water because of its antimicrobial and consumer-friendly properties.

Conclusion

Lettuces purchased from Kampar are safe to be consumed. Vinegar is effective to reduce the growth of *E. coli O157:H7* on RTE vegetables after a period. Nevertheless, various parameters including pH of vinegar shall be studied to determine its optimum effectiveness towards eliminating bacteria.

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0-7

Raman-based spectroscopic techniques for *Leptospira* DNA detection

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Background

Raman spectroscopy (RS) and surface enhanced Raman spectroscopy (SERS) are both promising techniques for biomolecule detection. In this article, we present our preliminary findings in the utilization of these techniques for *Leptospira* deoxyribonucleic acid (DNA) detection. A functional layer formed via a chemical organized monolayer (SAM) was used to immobilize DNA probe (pDNA) onto the metallic Raman layer/substrate. Successful hybridization between pDNA to the complementary DNA (cDNA) sequence was verified via vibrational features in Raman spectrum. The finding shows the potential use of Raman-based techniques as alternative diagnostic tools for leptospirosis.

Methodology

Two genes of *Leptospira*, i.e., *secY* and *lipL32*, were used. The following are the oligonucleotide sequences:

- SecY cDNA: 5'-TTT GAA GGG CAG GAA CAA G-3'
- LipL32 cDNA: 5'-TTG TTT CCA TCG ACT AAA CCG TC-3'
- SecY pDNA: 5'-/5AmMC6/CTT GTT CCT GCC CTT CAAA-3'
- LipL32 pDNA: 5'-/5AmMC6/GAC GGT TTA GTC GAT GGA AAC ΔΔ.3'

SecY gene is a housekeeping gene whereas lipL32 gene is a marker for pathogenic gene. The pDNAs were terminated with amine for rapid attachment to metallic Raman layer/substrate. A 50 nm gold (Au) thin-film and 80 nm in size bi-metallic gold- silver nanoparticles (Au-Ag NPs) were the metallic Raman layer/substrate for RS and SERS measurements, respectively. The Au thin-film was deposited onto a fused silicate layer by plasma sputtering, while the colloidal Au-Ag NPs were embedded onto the polymer-based photonic crystal substrate via polyethylenimine (PEI) adhesive. The metallic surfaces were SAM-modified; (1) for Au thin-film, 3-mercaptopropionic acid (MPA) and 1-ethyl-3-(3-dimethyl-aminopropyl) carbodiimide hydrochloride/ N-hydroxyl succinimide (EDC/NHS) were used to achieve thiolateand aminate-modified surfaces, (2) for Au- Ag NPs, the 4aminothiophenol (4-ATP) was used to obtain the amine tail on the surface. After the functionalization process, the metallic Raman layer/ substrate was incubated with 1 µM pDNA and cDNA in sequence. Raman spectrometer with 633 nm laser was used to investigate the vibration spectra from the samples.

Results and Discussion

Vibrational Raman peaks which correspond to chemical interactions and hybridization between pDNA and cDNA were observed from both RS and SERS measurements. A clear difference in Raman response can be seen on different surfaces. For Au thin-film, Raman peaks of phosphate backbone (PO $^-$) and adenine- (A), guanine- (G), thymine- (T) and cytosine- (C) nucleobases were observed to verify the hybridization of 1 nM secY cDNA. For Au-Ag NPs, the signal-tonoise ratio of SERS peaks after 1 μ M lipL32 cDNA hybridization is more prominent than the RS peaks. The characteristic of SERS intensity decreases with the presence of cDNA after hybridization.

Conclusion

The vibrational Raman spectra from *Leptospira* DNA was successfully detected via Raman spectrometer via RS and SERS techniques. SERS exhibits more intense DNA hybridization peaks in comparison to RS, thus offering better detection sensitivity.

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0-10

Performance evaluation of nested polymerase chain reaction (nPCR), light microscopy and *Plasmodium falciparum* histidine rich protein 2 rapid diagnostic test (*pfhrp*-2 RDT) in the detection of falciparum malaria in Akure, Nigeria

falciparum malaria in Akure, Nigeria
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Background

Malaria remains a serious public health problem worldwide. In order to ensure early and accurate malaria diagnosis, the World Health Organization recommended confirmatory parasitological diagnosis of malaria by microscopy and malaria Rapid diagnostic test (RDT) prior antimalarial administration and treatment. This study was designed to evaluate the performance of nested polymerase chain reaction (nPCR), light microscopy and *Plasmodium falciparum* histidine rich protein 2 rapid diagnostic test (pfhrp-2 RDT) in the detection of falciparum malaria in Akure, Nigeria.

Methodology

A cross-sectional and hospital-based study involving 601 febrile volunteered participants was conducted in Akure, Nigeria. Approximately 2-3 mL venous blood sample was obtained from each study participant for standard parasitological confirmation by microscopy and RDT. Thick and thin films were prepared and viewed under the light microscope for parasite density quantification and species identification, respectively. Dry blood spot (DBS) samples were prepared on 3MM Whatman filter paper for molecular analysis through nested polymerase chain reaction (nPCR).

Results and discussion

The overall prevalence by microscopy, RDT and nPCR were 64.89% (390/601), 65.7% (395/601) and 67.39% (405/601), respectively. Obviously, the performance efficacy of microscopy was significantly higher than RDT considering PCR as the gold standard. The estimates of sensitivity, specificity, Positive Predictive Value (PPV), Negative Predictive Value (NPV), accuracy, Youden's j index of microscopy and RDT were 96.30, 100.00, 100.00, 92.89, 97.50, 0.963 and 95.06, 94.90,

97.47, 90.29, 95.01, 0.899, respectively. Similarly, RDT recorded higher false negativity compared microscopy (4.94% vs 3.70%). A near perfect agreement was reported between microscopy and PCR, and between RDT and PCR with Cohen's kappa value (k) of 0.94 and 0.88, respectively.

Conclusions

This study revealed that RDT and microscopy continue to remain highly efficacious, though lower than PCR. Thus, while RDT continues to complement microscopy as the gold standard in high malaria-endemic settings, the application of PCR for constant evaluation and monitoring of RDT and microscopy is highly imperative to inform appropriate policy on malaria diagnosis and interventions.

0-11

Association of single nucleotide polymorphisms on ironregulating genes with iron metabolising parameters

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Background

Anaemia is a condition with haemoglobin concentration below the established cut-off level of 12 g/dL and insufficient to meet individual's physiological demands of oxygen. This condition has become a worldwide public health problem. Other than gender, family history, physiological condition, dietary intake, genetic factors contribute to the development of anaemia to a significant extent. In Malaysia, the prevalence of anaemia was up to 13.8% and more than half of the cases were due to iron deficient. Thus, this study is to elucidate the underlying factors in leading to anaemia including the association of SNPs on iron- regulating genes to iron-metabolising parameters and its predisposition to anaemia.

Methodology

A total of 183 subjects aged between 18 to 35 were recruited at Universiti Tunku Abdul Rahman with informed consent. Anthropometric measurement and haemoglobin (Hb) level were measured. Demographic data, family history and current physiological condition were self-declared by respondents through a questionnaire. A volume of 6 mL of venous blood was taken for DNA extraction while the isolated plasma and serum were used for detection of hepcidin and serum iron concentration. Eight genetic variants from 4 genes (rs10414846, rs10421768, rs855791, rs4820268, rs1799852, rs12769, rs1799945 and rs1800562) were genotyped using tetra primer- ARMS PCR. Statistical analysis was done using SPSS version 22.

Results and Discussion

The anaemia prevalence rate in this study population was found to be 14.75% (27/183). Women were found to have lower haemoglobin, hepcidin and serum iron level compared to men. Individuals currently menstruating or having the experience of menorrhagia presented with lower haemoglobin, hepcidin as well as serum iron. A lower hepcidin level is necessary to increase iron absorption to replenish iron loss through menstruation. Higher body mass index (BMI) was associated significantly with higher haemoglobin level, probably due to higher dietary iron intake among overweight and obese individuals. Individuals with a family history of anaemia presented with lower haemoglobin levels. The genetic variant of TF gene (rs12769) was associated with higher hepcidin and serum iron level while homozygous A allele in rs10421768 was associated with higher serum iron level. Other genetic variants were found to have no association with the iron-metabolising parameters that were analysed.

Conclusion

Anaemia is a multifactorial disease which could be due to gender, dietary intake, physiological condition as well as genetic factors. SNP in *TF* and *HAMP* gene modulates the hepcidin concentration as well as serum iron level. Findings can be further concluded in a larger sample size.

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0-12

Stable expression of anti-BmR1 IgG4 antibody

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Background

Lymphatic filariasis is a parasitic disease and one of the most disabling in the world. Thus, WHO called for elimination of this disease as a public-health problem in 1997. Diagnostic kits were made and distributed worldwide to address this problem, and one of the kits is the Brugia Rapid test (Reszon Diagnostic International Sdn. Bhd., Malaysia) that uses recombinant Brugia malayi antigen BmR1 which is highly specific and sensitive for antifilarial igG4 antibodies in patients with brugian filariasis. In the manufacturing of rapid tests, quality control (QC) is necessary to ascertain the reactivity and specificity of the tests [1]. An anti-BmR1-lgG4 antibody which was synthesized and verified as a QC reagent was produced through transient expression [2]. However, transient expression has its downsides of being laborious, having low productivity and high cost in the long term as compared to stable expression which requires much lower amount of plasmid DNA and act as a renewable source of expressing cells. Thus, this study investigates the conversion of transient to stable expression of the antibody.

Methodology

Stable cell lines were generated using Flp-In system and Flp-In 293 cell line (Thermo Fisher). DNA sequence of anti-BmR1 lgG4 was cloned into pcDNA5/FRT, which was then co- transfected with pOg44 into Flp-In 293 cell line using Lipofectamine 3000 (Thermo Fisher). True transfectants were selected based on Hygromycin B resistance and expanded to make cell stocks and working cells for antibody expression. Protein A affinity chromatography was used to purify the supernatant, which contained the protein, after the cells had been harvested. The respective recombinant antibodies were analysed qualitatively using

Western Immunoblot, as well as quantitively at 280nm using a nanodrop spectrophotometer.

Results and Discussion

Western immunoblot using anti-BmR1 IgG4 recombinant antibodies showed bands at the approximate molecular masses at 69.3kDa [2]. In 10 days, 1cm^2 of cell culture surface area (approximately 1.5×10^5 of cells) expresses up to 0.776µg of antibody. Transient or stable expression systems should be chosen based on throughput, the number and quality of resources required, and turnaround time.[3]. Stable cells are favoured for large-scale protein production because they offer high yields and reliable quality of no batch-to-batch variation. Site-specific recombination has the ability to consistently produce stable cells with high yields. The total surface area of a planar vessel can be increased to easily increase the stable cell pool.

Conclusion

Although transient system may be apt for initial studies, mass production of antibody would require a stable expression system. The

stably expressed antibody functions similarly to its transiently expressed counterpart. The Flp-In system is able to stably express the anti-BmR1 IgG4 antibody consistently and sustainably in an infinite manner, saving cost long-term.

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0-13

The role of Pp4R1 in T leukemic cell's survival

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Background

Reversible phosphorylation is one of the critical post-translational cell events, controlled by protein kinases and protein phosphatases. Protein kinases are involved in many human diseases including cancers. Protein phosphatases regulate cellular events by reversing protein kinases activity. The serine/threonine protein phosphatase 4 (PP4) is an essential protein for nucleation, growth, and stabilization of microtubules in centrosomes/spindle bodies during cell division and survival of T cells. It consists of catalytic subunit (PP4c) interacting with four different regulatory subunits (PP4R1, PP4R2, PP4R3 and PP4R4). Previous studies showed that PP4c has an important tumour suppressor function and plays crucial role in the control of cell fate in leukemic T-cells and untransformed human peripheral blood T-cells. The present study investigates the role of the protein phosphatase 4 regulatory subunit 1 (PP4R1) in leukaemia.

Methodology

Two different leukemic T cell lines Jurkat and CEM-C7 were transfected with two different PP4R1 specific siRNAs to study the effects of PP4R1 down-regulation on cell survival or with a plasmid encoding PP4R1 (pcDNA3.1-PP4R1) to investigate the effects of PP4R1 up-regulation.

Results and Discussion

Western blot analysis confirmed both down-regulation and overexpression of PP4R1. Reduction of PP4R1 protein levels was associated with an increase in the number of both total and viable cells. Increased levels of PP4R1 led to a significant decrease in the total and viable cell number, increase in basal apoptosis and cell cycle arrest in G1.

Conclusion

Overall, the results show that together with PP4c, PP4R1 regulates cell survival and growth of Jurkat and CEM-C7 leukemic T cells and propose that it could play an important role in maintaining the balance between cancer cell survival and death and might clarify the distinct pathological mechanism of leukaemia.

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0-14

X-chromosome wide association in Thai SLE patients

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Background

A previous study has shown that the genetic components especially in X chromosome contributed to female-biased found in Systematic Lupus Erythematosus (SLE) patients [1]. A copy of X chromosome is relatively correlated with SLE development [2]. Due to the global hypomethylation or defective in X-Chromosome lnactivation (XCI), [2] the studies found that 23% of genes are able to escape from this mechanism leading to X-linked gene over-expression found in SLE [3]. Genetic variations are deviated in different populations and may influence the incidence and mortality rate. Therefore, our study aims to investigate a global variant on X chromosome in Thai population. This might help to address the high female to male incidence ratio found among SLE patients.

Methodology

All procedures are approved by the ethics committee at Faculty of Medicine, Chulalongkorn University (EC. 590223). Using Asian Screening Array results from previous study [4], We conducted genomewide association study (GWAS) on X chromosome from 892 SLE samples compared with healthy controls data (n = 1,683). Low quality samples and bad SNP markers were excluded by quality control processes (QC). Inflation factors after QC process are 1.008. Next, haplotype estimating was completed by SHAPEIT to pre-phasing the SNPs before imputation step by IMPUTE v2.3.2 software using -chrX function. SNPs association analysis were analyzed using plink v1.9 with logistic correlation approach.

Results and Discussion

From our analysis, we characterized several known independent susceptibility SNPs in Thai SLE patients which include rs1059702 (G>A) (OR = 0.63, p-value = 9.30x10⁻⁹) located at *IRAK1- MECP2- TMEM187* region. Interestingly, the risk loci rs777448097 in *TLR7* (p-value 1.36 x 10^{-5} , OR = 0.77) were found in Thai population while it was absent in Hong Kong and Central Chinese background [1]. Furthermore, we found novel suggestive signal (rs14229594, C>T) on *GAB3-CTAG1A loci* (OR = 0.71, P = 2.14x10⁵). This variant showed higher effect size in female SLE patients and located within the enhancer region of CD14+ Monocytes (ENCODE database).

Conclusion

In the present study, we report a number of known SLE susceptible loci, specifically in the Thai population. Although the suggestive novel risk loci on *GAB3-CTAG1A* regions have been identified, further studies are required to validate this result. Our finding may help address the female biased phenomenon associated with SLE in the Thai population.

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0-15

The effect of mesenchymal stem cells-mediated macrophages activation on breast cancer progression

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Background

Breast cancer (BC) is the second most leading cause of cancer-related death among women worldwide and represent about 11.7% of all new diagnosed cancer cases in 2020 [1]. High population of tumor-associated macrophages (TAM) in tumor microenvironment (TME) which display M2-type macrophages promotes BC metastasis and correlates with poor clinical prognosis in BC patients. Reprogramming TAM into M1-type macrophages that capable of killing tumor cells has emerged as a favorable therapeutic target in BC. Mesenchymal stem cells (MSC) can stimulate immunomodulatory changes and skew naïve macrophages (M0) to M1-type macrophages [2]. Therefore, this study attempted to evaluate MSC-macrophages crosstalk on BC progression.

Methodology

THP-1 cell was incubated with PMA for 48 hours to differentiate into M0. M0 were co-cultured with MSC for 30 hours. Conditioned medium (CM) was collected to evaluate M1- and M2-type macrophage related cytokine while total RNA was extracted from co-cultured cells and then synthesised into cDNA to determine IRF-4 and IRF-5 mRNA gene expression using RT- PCR. Finally, MDA-MB-231 cells were cultured with CM of respective treatments for 24, 48 and 72 h to evaluate their cell viability using MTT assay.

Results and Discussion

Results showed that UC-MSC significantly increased IRF-4 expression in M0 and M0+LPS compared to M0 alone but no effect on IRF-5. In addition, UC-MSC did not influence the secretion of IL-10 and TNF-α in the supernatant of M0 and M0+LPS. Moreover, CM from M0, M0+UC-MSC, M0+LPS and M0+LPS-UC-MSC significantly reduced the proliferation of MDA-MB-231 cells at 72 h compared with 24 h treatment respectively. The results indicated that UC-MSC induced the polarisation of M0 into M2-type phenotype as IRF-4 regulates M2-type macrophage polarisation. M0 secretory products suppressed ER- breast cancer cells (MDA-MB-231) as observed in a previous study (3). UC-MSC and LPS may also stimulate the M0 to secrete a secretome that contains anti-cancer properties that hinders the proliferation of triple negative BC cells. This finding is consistent with a prior study that showed umbilical cord matrix derived MSC-CM inhibited the cell viability of BC cells.

Conclusion

Our findings demonstrate that UC-MSC may polarise naïve macrophages into M2-type macrophages. UC-MSC and LPS work in synergy in stimulating M0 macrophages to secrete anti-cancer products that suppress the growth of triple negative breast cancer cells.

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0-16

Therapeutic properties of Malaysian stingless bee pollen and its protective effect against DNA damage

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Background

Reactive Oxygen Species (ROS) can disturb cellular metabolism and damage cellular biomolecules, which could lead to DNA damage. Natural compounds including bee-collected pollen contain nutrient antioxidants that have therapeutic properties and protective effects against ROS. Bee pollen is known as a complete food since the food energy produced is relatively high, thus it serves as a source of nutrients for adult bees and larvae [1]. Studies demonstrated that 70% of bee pollen compositions are biologically active and exhibit numerous benefits including nutrition, cardioprotection, hepatoprotection, immunostimulant, antioxidation, anticarcinogen, antibacterial, antiosteoporosis, antiprostatitis, anti-anaemia, anti-ageing, and anti-inflammatory [2, 3]. This study aims to investigate the protective effect of stingless bee pollen against DNA damage.

Methodology

Bee pollen ethanolic extracts (BPEs) were prepared from three stingless bee species commonly domesticated in Malaysia: i.e., *Tetrigona apicalis, Heterotrigona itama*, and *Geniotrigona thoracica*. The methodologies used in this study were spectrophotometric method for total phenolic/flavonoid content and antioxidant activities, HPLC and GC-MS chromatographic techniques for phenolic compounds identification, trypan blue exclusion assay for antiproliferation test, and comet assay for DNA damage activities. MCF-7, MCF-10A and HT29 cell lines were used in this study.

Results and Discussion

In antioxidant assay, the result showed that BPE from Malaysian stingless bee species contained different phenolic and flavonoid contents, with each extract possessing different amounts of antioxidant activity. G. thoracica BPE possessed the highest capacity to neutralize DPPH radicals compared to T. apicalis and H. itama. HPLC and GC-MS analysis detected various polyphenol compounds and chemical groups in each species. The chemical compounds found in BPE have lots of biological activities with antioxidative potential to be explored as health supplements and medical treatment. In antiproliferation assay, G. thoracica exhibited the highest therapeutic index (Tl=3.12), with the EC₅₀ of 0.5 mg/mL in HT-29 cells. With the therapeutic potential, stingless bee pollen may provide a preventive function against ROS and the development of many lifestyle diseases. Analysis of DNA damage activity at 24 h of G. thoracica treatment showed a significant decrease in H₂O₂-induced DNA damage compared to the untreated cells (63.82% + 2.46 vs. 90.86% + 0.68) (p<0.01). Similarly, a significant reduction was also seen with the supplementation of caffeic acid (49.05% + 4.23) and quercetin (43.98% + 3.77) (p<0.01). More significant reduction of DNA damage was observed at 72 h of *G. thoracica BPE* (20.49% + 0.73), caffeic acid (5.65% + 0.35), and quercetin treatment (7.58% + 0.32) (p<0.01) on H_2O_2 -exposed HT-29 cells. Due to its strong antioxidant activity, *G. thoracica* BPE gave a protective effect against H_2O_2 -induced DNA damage.

Conclusion

Results of the present study discovered that Malaysian stingless bees, particularly G. thoracica species have a genoprotective effect that can protect cells from H_2O_2 -induced oxidative DNA damage. Even though the exact mechanism concerning the protective effects of BPE is still unconfirmed, the bioactive phytochemicals and antioxidant capacity are believed to be responsible for the suppression of oxidative DNA damage and may confer protection from genetic instability.

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D-18

Acute and sub-chronic toxicological evaluation of probiotic strain *Lactobacillus rhamnosus* GG in Sprague Dawley rats

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Background

Lactic acid bacteria are one of the major groups of gastrointestinal bacteria in healthy humans. A strong history of commensal status and long-term usage without any obvious adverse effects has classified them as 'Generally Recognized as Safe'. In recent years, bacteriotherapy using probiotics as therapeutic agents has shown to be a promising emerging method for evasion of infectious diseases. *Lactobacillus* spp. as probiotics has been questioned for its safety due to recently reported unexpected reactions. These include *Lactobacillus*-associated systemic infections, metabolic reactions, immune disorders and gene transfer etc.]. Among the probiotics, opinions proposed by several organizations between countries remain still contradictory. Although *Lactobacillus* spp. as probiotics have been classified safe, *Lactobacillus rhamnosus* strain GG requires further surveillance with additional studies.

Methodology

Acute and sub-chronic toxicity studies of probiotic strain *Lactobacillus rhamnosus* GG in Sprague-Dawley rats (180 \pm 20 g) as per OECD TG 423 and 407, respectively. Acute oral toxicity study was conducted by the administration of 1 \times 108, 1 \times 109 and 1 \times 1010 cfu/ ml of *L. rhamnosus* (n = 3/ group). The animals were observed for their behavioral, neurological and autonomic functions continuously for 24 h and monitored 14 days thereafter for mortality. Sub-chronic toxicity study was conducted by the administration of 1 \times 106, 1 \times 107 and 1 \times 108 cfu/ ml of *L. rhamnosus* once daily for a 28 days period (n = 12/ group). During the study period, the animals were monitored for changes in body weight and behavioral functions. The blood sample

was collected on day 14 and 28, used for biochemical and hematological parameters analysis. At the end of study the animals were sacrificed and pivotal organs were collected and used for histopathological analysis.

Results and Discussion

In acute toxicity studies, *L. rhamnosus* did not show any changes in behavioral, neurological and autonomic functions, and mortality. In sub-chronic study, *L. rhamnosus* did not show any significant changes in body weight gain, organ weight analysis and behavioral functions. *L. rhamnosus* at 1×10^6 cfu/ml did not show any significant changes in biochemical and hematological parameters when compared with the control. Whereas *L. rhamnosus* at 1×10^7 and 1×10^8 cfu/ml showed dose-depended changes in elevated aminotransferase levels when compared with the control. In histopathological analysis, mild degeneration of hepatic cells and nephrons were observed in *L. rhamnosus* at 1×10^8 cfu/ml administered group when compared with the control.

Conclusion

The probiotic *L. rhamnosus* GG did not show any significant toxic effect at 1×10^6 cfu/ml and exhibited mild-to-moderate toxic effects at the dese levels of 1×10^7 cfu/ml and 1×10^8 cfu/ml. Pre-clinical toxicity testing helps to calculate "No Observed Adverse Effect Level [NOAEL]".

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0-19

In-depth investigation of microRNA methylome signature in colorectal cancer

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Background

Colorectal cancer (CRC) is one of the top causes of cancer-related deaths worldwide. Despite substantial breakthroughs in diagnostic services and patient care, various gaps remain to be filled, ranging from early detection to the identification of prognostic indicators, effective therapy for metastatic illness, and the implementation of personalised treatment methods. MicroRNAs, which are small noncoding RNAs, are unregulated in CRC and play an important role in its onset and progression. Despite this, microRNA research has traditionally relied on expression levels to assess biological importance [1]. The precise mechanism underlying microRNA dysregulation in cancer has yet to be determined, however multiple studies have shown that epigenetic mechanisms, notably DNA methylation, play essential roles in the regulation of microRNA production. Thus, we

intend to investigate the miRNA methylome patterns and explore their roles CRC.

Methodology

Fifty-four pairs (n=108) of CRC and the respective adjacent normal tissues were collected from the UKM Medical Center, Malaysia. Methylation profiling was performed using the Infinium Human Methylation 450K beadchip, which covers 485,577 CpG dinucleotide sites distributed over the whole genome. The raw IDAT files were exported from the scanner, and quality control was performed using Genome Studio and were further analysed using the ChAMP R package.

Results and Discussion

We obtained 933 miRNA probes for the downstream analysis. These probes were further classified as hypermethylated or hypomethylated based on the absolute average β value difference ($\Delta\beta$) at \geq 0.2 between CRC and normal adjacent tissues. A total of 230 probes were identified. Distribution of hypomethylated miRNAs with respect to genomic regions revealed more than half of the miRNA probes were located in the TSS1500 (n = 130; 58%), followed by TSS200 (n=55; 24%) and body (n= 40; 18%). On the other hand, the distribution of hypermethylated miRNA probes is almost equal in TSS200 (n=3; 60%) and TSS1500 (n=2; 40%). Meanwhile, categorization based on CpG islands (CGIs), 225 probes were hypomethylated and five probes were hypermethylated. The five hypermethylated miRNA probes were MIR1180_cg24553547, MIR124-3 cg02650317, MIR34B cg22879515, MIR124-3 cg15699267 MIR137_cg22333214. Among these, MIR1180_cg24553547 showed the most significant differentially hypermethylated delta β-value. Although atypical methylation status has been documented in tumour samples, the methylation profile of miR-1180, which is located on chromosome 17, has yet to be thoroughly explained in CRC. Prior research demonstrated a good association between hypermethylation of the miR-1180 gene and downregulation of its expression in tumour tissue when looking at its expression level in malignancies. Unlike other miRNAs, miR-1180 functions as both an oncogene and a tumour suppressor. Tan et al. [2] and Zhou et al. [3] previously demonstrated the carcinogenic activity of this miRNA in hepatocellular carcinoma by directly targeting both OTUD7B and TNIP2 to drive cell proliferation and apoptosis resistance.

Conclusion

Our findings imply the existence of new pathways involving methylation alterations impacting miRNA genes that control CRC development. The new knowledge from this study can be utilized for personalized health diagnostics, disease prediction, and monitoring of treatment.

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0-20

Identification of LRRC17 as colonic fibroblast activation marker and its potential role in colorectal cancer progression

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Background

Stromal fibroblast is linked with poor prognosis of colorectal cancer (CRC) [1]. Crosstalk between cancer-associated fibroblasts (CAFs) and cancer cells facilitated by growth factors drives the malignant

progression. CAFs can be identified using conventional markers such as alpha-smooth muscle actin (α -SMA) although their expression can be heterogenous. Leucine Rich Repeat Containing 17 (LRRC17) is proposed to be an emerging biomarker to specifically identify activated fibroblasts and CAFs. This study was aimed to verify LRRC17 expression in different colonic fibroblasts and further characterize LRRC17 in CRC. This may provide insights on the underlying mechanisms and verify its potential as a specific marker for future prognostic purposes [2].

Methodology

LRRC17 and α -SMA expressions were analyzed using immunofluorescence staining on CCD-112CoN (MF cell line) and primary fibroblast lines derived from colorectal tumor and normal adjacent colon tissues (denoted as normal fibroblasts). SW620 (CRC epithelial cell line) was used as control. The cells were cultured under different conditions; a) serum-free medium (DMEM alone), b) DMEM + 10% fetal bovine serum (complete medium), c) conditioned medium (CM) of SW620, and d) DMEM + transforming growth factor β 1 (TGF- β 1). Fluorescence intensity was assessed via ImageJ 1.53k. Treated cells proliferation was assessed via MTT assay. All experiments were conducted in duplicates.

Results and Discussion

Positive LRRC17 and α -SMA expressions were observed in all fibroblast groups and CCD-112CoN except SW620. However, different co-localization patterns were shown between the proteins in treated cells. Both proteins expression was greater in complete medium and TGF- β 1 as compared to serum free group, indicating more activated state of treated fibroblasts mimicking CAF property. Interestingly, higher LRCC17 expression whereas lower α -SMA expression in CM SW620 for primary cell lines. This may highlight the cancer secretome effects which trans-differentiate fibroblasts or change their molecular property. Different expression patterns seen in the present study may indicate different pathways involved in the regulation of these markers, suggesting further study. From MTT assay, highest proliferation was found in fibroblasts treated with complete medium. It was proposed that growth factors in serum support the cell growth [3].

Conclusion

LRRC17 can potentially serve as biomarker to identify and characterize CAFs of CRC.

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0-21

Effect of polyphenolic-rich fraction of cornsilk (*Stigma maydis*) in streptozotocin-induced diabetic rats

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Background

Diabetes is a chronic condition that arises when the pancreas does not create enough insulin or when the insulin produced is not used efficiently by the body. Hyperglycemia, or elevated blood sugar, is a frequent complication of untreated diabetes and can cause catastrophic harm to several of the body's systems, including the neurons and blood vessels. Current available drug such as metformin which is widely used to treat hyperglycemia have some side effects such as diarrhea, vomiting and lactic acidosis. Currently, there is a keen interest in studying the benefits of food waste, hence researchers started to study the benefits of natural product food waste in treating many pathological conditions. Therefore, the aim of this research is to study the effects of polyphenolic-rich fraction of cornsilk in reducing the fasting blood glucose level in diabetic-induced rats.

Methodology

The effect of polyphenolic-rich fraction (PRF) of corn silk (CS) extract were studied on the normal cell line Human Umbilical Vein Endothelial Cells (HUVEC) and Streptozotocin-induced diabetic rats Streptozotocin-induceddiabetic rats.

Results and Discussion

The proliferation assay result of PRF of CS obtained shown significant percentage of cell viability when compared to metformin (p<0.05). The effects of PRF of CS extracts in fasting blood glucose level shown no significant different between treated and untreated group (p>0.05), might be due to the 14-day duration of this study. The effects of PRF of CS on the body weight of rats also shown no significant different when compared between treated and untreated group (p>0.05).

Conclusion

However, the histopathological study shows promising effect of PRF of CS in treating diabetes when compared to untreated diabetic group by assessing the morphology of the glomerulus, tubules and Bowman space in kidney and the morphology of centrilobular area, hepatocytes and sinusoidal space in the liver. The long-term impact of PRF on CS in diabetic models has to be studied further by repeating the research study over a longer time period.

0-22

Phytochemical, anti-microbial activity and anti-proliferation test against human cancer-origin cell lines using water extracts of *Momordica cochinchinensis* (Gac fruit)

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Background

Momordica cochinchinensis (Gac fruit) is a seasonal tropical fruit that is widely used in Southeast Asian countries [1]. However, it's not fully exploited in Malaysia. The major component of this fruit that has been focused on is the aril, however the pulp, peel and seed are often discarded. Therefore, this study aims to scientifically validate the presence of phytochemicals composition, determine the antioxidant, antimicrobial and anti-proliferative activities of different parts (aril, pulp and seed) of Gac fruit grown in Malaysia.

Methodology

The fruit (aril, pulp and seed) was macerated and extracted using water following the traditional preparation approach. A toxicity test using brine shrimp (BSLT), phytochemical tests, antioxidant, antimicrobial, antifungal and cytotoxicity cell-based assays were conducted.

Results and Discussion

The crude extract showed 50% mortality in brine shrimps after 24 hours and its LC_{50} -value was considered moderate toxic. Qualitative phytochemical analysis revealed the presence of alkaloids, flavonoids,

saponins, volatile oils, glycosides and tannins. The highest scavenging activity of water extract from the pulp with 30.1 mg GAE/g FW and 0.012 mg GAE/g FW in both DPPH and FRAP, respectively. The pulp water showed high levels of total phenolic and flavonoid content of 0.0215 mg GAE/g FW and 0.083 mg QE/g FW, respectively. Moreover, pulp water extract displayed strong antimicrobial activity with MIC values ranging from 5-20 mg/ml and MBC values of 10-20 mg/ml on certain microbial strains. Cancer origin cells MCF7, HepG2, A549, HCT116 and HT29 were found to be more susceptible with the treatment of aril and pulp water extracts with the LC50 at 15.41, 5.87, 22.89, 1.10 and 0.03 µg/ml while pulp at higher values such as 269.40, 328.46, 34.90, 68.07 and 21.44 µg/ml at 72 hours post-treatment compared to other tested cell lines.

Conclusion

The results concluded that Gac fruit from crude extracts revealed chemical constituents (alkaloids, flavonoids, saponins, volatile oils, glycosides and tannins) contributing to their antioxidant and antimicrobial activity against *E.coli, P.euroginosa, B.cereus and S.flexneri.* Further studies on the mechanism pathways need to be explored since Gac fruit showed antiproliferative activity against breast, liver, lung and two colon cancer cell lines.

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0-23

Conjugates between P. marcocarpa aqueous extract and TiO₂ exhibited a synergistic antimicrobial effect

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Background

Phaleria marcocarpa (P. marcocarpa), also known as Mahkota Dewa is a flowering plant of Thymelaeceae family. Originated from Indonesia, it thrives in tropical climate countries such as Malaysia. It has been reported to possess anti-diabetic and anti-bacterial properties and is often brewed as tea for consumption. However, the antimicrobial activities of P. marcocarpa are proven to be weak and limited to limited strains of microbes. Titanium dioxide nanoparticles (TiO₂ NPs) are semiconducting transition metals with strong antimicrobial activities. P. marcocarpa aqueous extract was conjugated with TiO₂ NPs in this study to determine the synergistic antimicrobial effect of the resultant conjugate against Escherichia coli (E. coli) and Staphylococcus aureus (S. aureus).

Methodology

The aqueous P. *marcocarpa* fruit extract was extracted through maceration for 24 hours. The extract was filtered through Whatman® filter paper No 1 and freeze dried. TiO $_2$ NPs were synthesized through the sol-gel method. The conjugation between the aqueous extract and TiO $_2$ NPs were carried out at various temperatures (30°C, 45°C and 60°C). The resultant conjugate was characterized through FTIR. The antimicrobial activity test was carried out through MTT assay against *E. coli* (ATCC 25922) and *S. aureus* (ATCC 6538). The assay was carried out using a 96-well sterile microtiter plate. The concentration of extract, TiO $_2$ NPs and resultant conjugate used were from 125 to 15.6 μ g/mL.

Results and Discussion

The FTIR results showed difference between the TiO₂ NPs and resultant conjugate around the fingerprint region. Bands at 3305, 2129, 1637, 1538 and 1370 cm⁻¹ have been shown by pure TiO₂ NPs, while bands at 3305, 2144, 1637 cm⁻¹ have been shown by the extract. Additional band at 1045 cm⁻¹ was shown by the resultant conjugate while additional band at 792 cm⁻¹ was observed for the conjugate prepared at 60°C. The 1045 cm⁻¹ band represented anhydride and alcohol stretching while the 792 cm⁻¹ band represented the bending of alkene. From the antimicrobial test against E. coli and S. aureus, the conjugates prepared at 30°C, 45°C, and 60°C has minimum inhibitory concentration (MIC) of 15.6 µg/mL. However, the conjugate at 45° C showed lowest CFU/mL of 3.5×10^{6} for E. coli and 9.6×10^{6} for S. aureus when compared to TiO₂ NPs and other conjugates (30°C and 60°C) at the same concentration. Thus, it was proven that 45°C is the optimum temperature for TiO2 extract conjugation in view of growth inhibition for E. coli and S. aureus as it showed lower CFU/mL as compared to the pure TiO₂ NPs.

Conclusion

In conclusion, the MTT assay was effectively used at evaluating the antimicrobial activity. The resultant conjugate showed the most remarkable synergistic antimicrobial effect of MIC at 15.6 μ g/mL and 45°C for both E. *coli* and S. *aureus*.

0-24

Cytotoxicity, proliferation and migration assessment of BHMC, the curcuminoid analogue on human liver cancer cells, HepG2

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Background

Over the years, natural bioactive compounds were acknowledged for their cytotoxic and anticancer capabilities. Curcumin is a bioactive compound derived from the rhizomes of Curcuma longa known to possess various pharmacological properties e.g., inhibiting cancer growth, exerting cytotoxic and inducing apoptosis [1]. Nonetheless, curcumin has some limitations that prevent it from reaching its full potential. Due to poor bioavailability, 2,6-bis(4-hydroxy-3-methoxy-benzylidene)cyclohexanone (BHMC), a curcuminoid analogue was synthesised via chemical structure modification by eliminating the unstable β -diketone moiety and converting it into a conjugated double bond while retaining the phenolic hydroxyl group to overcome the limitation by exerting greater cytotoxic, growth suppressive effects with more selective on cancer cells [2]. Therefore, it would be beneficial to look at the cytotoxicity along with the anti-proliferation and -migration effect of BHMC on HepG2 cells.

Methodology

The HepG2 and Hs27 cells were treated with various concentrations ranging from 0.781-50 μ M for 24 and 48hrs via MTT assay. Based on the IC50 value obtained, several concentrations were selected to be used in Trypan Blue Exclusion (TBE) assay, Migration assay and Hoechst/Propidium Iodide (PI) staining. Data were analysed using GraphPad Prism software.

Results and Discussion

BHMC was observed to be approximately 3-5 times more toxic to HepG2 compared to curcumin with IC50 values of 16.73 μM and

46.03µM at 24hrs while 4.77µM and 26.00µM at 48 hrs, respectively. However, BHMC was less toxic towards Hs27 cells with the IC₅₀ value of more than 30µM at 24 and 48 hrs. BHMC significantly reduced HepG2 by 30-60% at 24hrs and 70-80% after 48hrs at lower concentration than curcumin. In contrast, BHMC only reduced 20-30% of Hs27 at 24hrs and 40-60% at 48hrs. Curcumin reduced 40-60% of HepG2 for 24 and 48 hrs but only reduced 30% of Hs27 cells at similar concentrations and timepoint. Although BHMC and curcumin reduced the percentage of HepG2 cell viability in a concentration- and time-dependent manner, the anti-proliferative effect exerted by BHMC was greater compared to curcumin. BHMC also showed a cytotoxic selective effect on Hs27 especially at 48 hrs with less cell death at lower concentrations compared to HepG2. It is suggested that BHMC was cytotoxic selective towards Hs27 like its parental compound, curcumin. BHMC and curcumin suppress the migration of HepG2 as compared to controls. However, below the IC₅₀ value, BHMC exhibits a higher effect in suppressing HepG2 cell migration compared to curcumin. At lower concentrations, characteristics of apoptosis can be observed including cell shrinkage, membrane blebbing, chromatin condensation, DNA fragmentation and apoptotic hodies

Conclusion

BHMC mediates a greater cytotoxic effect in HepG2 by reducing the cell viability, modulating a higher percentage of cell death at lower concentrations compared to curcumin, suppressing migration, and inducing apoptosis compared to Hs27 while cytotoxic selective towards Hs27 cells at similar concentrations.

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0-25

Unraveling the tumour-regulatory role of miR-3934 in human breast cancer

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Background

MicroRNAs (miRNAs) are short, single-stranded, and non-coding RNAs that could regulate the expressions of multiple downstream cellular targets post-transcriptionally. By altering the expressions of tumour-promoting and tumour-suppressing targets, miRNAs could act as tumour-modulatory miRNAs in different human cancer. The preliminary findings from our previous work suggested that miR-3934 may be responsible in promoting tumour growth and migration in human breast cancer cell lines [1]. Therefore, this mechanistic study aimed to elucidate the tumour-regulatory role of miR-3934 in human breast cancer cells using *in vitro* cellular assays.

Methodology

Triple-negative human breast cancer cell, MDA-MB-231, was stably transfected with miRNA expression plasmids to produce clones that overexpressed miR-3934 mimic, miR-3934 inhibitor, or their respective scrambled control sequences. All transfected and un-transfected cancer cell types were subjected to cell viability, invasion, and

wound-healing assays to evaluate the tumour-regulatory role of miR-3934 in breast cancer cells.

Results and Discussion

Overexpression of miR-3934 significantly (p<0.05) promoted breast cancer cell proliferation and resistance to chemotherapeutic agent, cisplatin. Besides, miR-3934 also enhanced (p<0.05) the invasion and migration abilities of the MDA-MB-231 cell line as compared to untransfected control and clones that expressed scrambled control sequences. In contrast to clones that overexpressed miR-3934, clones that overexpressed miR-3934 inhibitor were shown (p<0.05) to have lower proliferation rate, higher sensitivity to killing by cisplatin, and have decreased invasion and migration abilities in vitro as compared to all other cell groups. There were no statistically significant differences in the cell proliferation, chemosensitivity to cisplatin, invasion, and migration rates between the untransfected control and clones that expressed the scrambled control sequence. The tumourpromoting role of miR-3934 observed in the current study is consistent with the findings observed in two other miR-3934 studies that involved lung cancer [2] and neuroblastoma [3], in which miR-3934 was reported to downregulate the expression of tumour suppressor protein such as TP53INP1 [2,3]. However, to date, the tumourmodulatory role and downstream target of miR-3934 in breast cancer are still unclear and under-reported. The current study findings suggested that miR-3934 has the potential to be utilised as a prognostic breast cancer biomarker and the introduction of a miR-3934 inhibitor may potentially inhibit breast cancer development. To further confirm the tumour-promoting role of miR-3934, transcriptomics and in vivo studies will be conducted to unveil the molecular mechanism of how miR-3934 promotes breast cancer development.

Conclusion

Overexpression of miR-3934 was shown to promote the proliferation, chemoresistance, invasion, and migration in the triple-negative MDA-MB-231 cell line while the miR-3934 inhibitor may potentially halt the tumourigenesis process in breast cancer.

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0-27

Selection of T-cell receptor (TCR) like antibody against Human Leukocyte Antigen A-2 (HLA-A2) for cervical cancer diagnostics

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Background

Cervical cancer is a leading cause of death among women. Persistent human papillomavirus (HPV) contributes to the carcinogenesis of cervical cancer, in which 70% of all cervical cancer cases are linked to HPV 16 and 18. The available HPV vaccines effectively prevent cervical cancer but do not treat existing cervical cancer. Besides, cervical

malignancy detection at early phases remains a challenge. Consequently, treatment begins at the late stages of the disease. Hence, new diagnostics and therapeutics methods are required to treat cervical cancer. In this study, a novel antibody class, known as T cell receptor (TCR)-like antibodies, specific to HPV 16 and 18 E7 oncopeptides, has been hypothesised to enhance cervical cancer diagnostic potentials. The dual functionality of the antibody in mimicking the function of T cell receptor (internal immunosurveillance) and simultaneously executes its antibody effector mechanisms to eliminate a disease effectively opens a new and promising pathway to target cervical cancer.

Methodology

Bioinformatics were used to select the target peptides specific to HLA-A2 of HPV 16 and 18 E7. Peptide-MHC (pMHC) complex was formed by refolding HPV peptides with HLA-A2 heavy chain and $\beta 2m$ light chain. Potential TCR-like antibodies were generated by panning the pMHC complexes against an antibody phage display library. The monoclones were analysed with monoclonal ELISA and comparative ELISA before sequence analysis.

Results and Discussion

The successful refolding of pMHC was analysed via ELISA with refolded β2m as positive control. A domain antibody library was used to select potential TCR-like antibodies specific to HPV 16 and 18 E7 pMHC by biopanning. Three rounds of biopanning were performed to get highly enriched antibody phages. Antibody phages from round three polyclonal panning were selected for monoclonal analysis because they have highest enrichment. For HPV 16 E7, 20 out of 92 clones were selected for monoclonal analysis and finally 9 clones were sent for sequencing. Meanwhile, for HPV 18 E7, 44 out of 92 clones were selected for monoclonal analysis and finally 17 clones were sent for sequencing. It was difficult to generate a novel group of TCR-like antibody until latest development in technologies like phage display techniques made it possible. The coat protein of a bacteriophage is fused with a protein/peptide and presented on a virion's surface in phage display technique. Antibody phage display technology has emerged as a famous approach for production of antibodies against various targets including cancer. TCR-like antibody is development based on distinctive roles of T (T cell receptor) and B cells (antibody). Generally, T cell receptor recognises the antigenic peptide presented by MHC complex on all nucleated cell while antibodies generated by B cells can differentiate soluble or membrane bounded antigens which will result in the elimination of illness due to its wider effector functions. Hence, dual functionality of TCR-like antibody in combining both humoral and cell mediated immunity in a single approach makes it an appropriate candidate for cervical cancer diagnosis.

Conclusion

Two out of 9 clones and 7 out of 17 clones for HPV 16 E7 and HPV 18 E7, respectively had proper sequence which were analysed by VBASE2 and IMGT/V databases. In the downstream process the proteins of positive antibody clones will be expressed and purified. The purified antibodies will be tested for their binding ability using ELISA and western blot. Finally, the antibodies will be tested in cervical cancer cell lines.

0-28

Development of T-cell receptor-like antibody against Human Leukocyte Antigen-A11 (HLA-A11) human papillomavirus (HPV) type 16 & 18 oncoprotein E7 for the diagnosis of cervical cancer

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Background

Cervical cancer is the fourth most common cancer in women worldwide. Human papillomavirus (HPV) is the causative agent of cervical cancers with HPV 16 and 18 being the predominant types causing more than 70% of all cervical cancer cases. Cervical cancer can be prevented through HPV vaccination. However, diagnosing cervical cancer at an early stage remains a challenge. Therefore, a new diagnostic tool is required. In this study, TCR-like antibodies which are a novel class of antibodies that are specific to E7 oncoproteins of HPV 16 and 18 have shown to have the ability to enhance the diagnosis of cervical cancer. The ability of TCR-like antibodies in sandwiching the humoral and cell-mediated immunity in a single approach has brought light to a new promising pathway for targeting cervical cancer.

Methodology

Target peptides which are specific to Human Leukocyte Antigen A-11 (HLA-A11) HPV 16 and 18 E7 oncoproteins were identified and selected by bioinformatics. Peptide-MHC (pMHC) complex was formed by refolding the target peptides with heavy chain (HLA-A11) and light chain (β 2m). Refolded pMHC were used to develop TCR-like antibodies via biopanning against the antibody phage display library. Positive monoclones obtained were sent for sequencing before being analyzed using VBASE2 and IMGTV databases.

Results and Discussion

The successful refolding of pMHC complexes for both HPV 16 E7 and HPV 18 E7 were analysed via ELISA using refolded β2m as the positive control. The refolded pMHC complex was subjected to three rounds of biopanning against antibody phage display library for selection. Phages from the third round of biopanning were selected for both HPV 16 E7 and HPV 18 E7based on the polyclonal ELISA results as they showed the highest enrichment compared to the first and second rounds. 94 single colonies for each HPV 16 E7 and 18 E7 were randomly selected from the phages' titration for monoclonal biopanning. 10 out of 94 potential monoclones for HPV 16 E7 and 37 out of 94 potential monoclones for HPV 18 E7 were subjected to comparative biopanning based on the monoclonal ELISA results with a value after normalization of \geq 0.3. A total of seven out of 10 potential monoclones for HPV 16 and 26 out of 37 potential monoclones for HPV 18 were sent for sequencing based on the monoclonal ELISA results with a value after normalization of ≥0.2. TCRlike antibodies are based on the distinct roles of T cell receptors that are involved in immunosurveillance and direct killing, and the B cell is involved in antibody production. TCR-like antibodies function by differentiating the antigenic peptide which is presented on the MHC molecules and establishing primary defence mechanisms of the antibody. Therefore, the duality of TCR-like antibodies' function makes it a great candidate for cervical cancer diagnosis.

Conclusion

In conclusion, one out of seven clones for HPV 16 and three out of 26 clones for HPV 18, respectively had proper sequences. In the downstream process, the proteins of the positive clones will be expressed and purified. Later, the purified protein will be evaluated for its binding capabilities using Western blot and ELISA. Finally, the antibodies will be evaluated on the cervical cancer cell lines.

0-29

High throughput molecular profiling of bacterial diversity in Johor mangroves

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Background

Biomedical research heavily relies on the discoveries of natural compounds and thus it is very important to safeguard nature to build resilience in biomedical research. Mangrove is the coastal ecosystem that has proven to harbour several beneficial microbial communities with a

wide range of therapeutic molecules such as antimicrobial agents, anticancer compounds and enzymes. Mangrove provides habitat for several organisms including bacteria that are adapted to the microclimate with high salinity and tidal variation [1]. Some of these bacteria include actinobacteria which form the major source of antibiotics. It is important to identify and discover new antibiotics as there is an increasing threat of resistance to currently available antibiotics and limited discovery of new antibiotics [2]. Mangrove biodiversity in rapidly developing parts of the globe is severely threatened due to the destruction of habitat and thus there is a high threat to the bacterial community. One of the major hurdles in studying mangrove soil bacteria is that it is tedious to culture them and hence difficult to get a complete profile of bacterial diversity by conventional methods. Thus, it is important to adopt high throughput molecular techniques to rapidly identify bacteria to accelerate conservation efforts [3].

Methodology

DNA was isolated from mangrove soils sediments from eight mangrove sites around Johor, Malaysia that are grouped based on their vulnerability status. The library was prepared using the multiplexing method with 24 barcodes to distinguish samples and sequenced for the 16S gene using Oxford Nanopore Technologies MinION sequencing. Base-calling of sequencing reads was performed by the Guppy programme and analysed using EPI2ME software. Quantitative PCR was performed to verify linear amplification.

Results and Discussion

We have successfully sequenced the bacterial 16S gene in mangrove soil samples from Pulau Kukup, Sungai Pulai, Sungai Melayu, Sungai Danga, Sungai Skudai, Sungai Johor, Sungai Sedili Besar and Sungai Sedili Kechil. We show that the barcoding would enable parallel sequencing of samples from all sites in the same run and amplicons showed linear amplification. We achieved over 6 million high-quality reads per run in less than 24 hours. We report a widespread difference in bacterial diversity between protected mangroves compared to mangroves that have been exposed to anthropogenic activities. This is the first study to show the bacterial community profile of Johor mangroves.

Conclusion

MinION sequencing is a quick and efficient way to evaluate bacterial diversity in environmental samples. The findings of this study would enable the development of bacterial community mapping as a repository to help in conservation/rehabilitation as well as monitor the impact of mangrove loss on bacterial diversity.

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O-30

Ranking of the *Mycobacterium tuberculosis* T-cell epitopes using tabulated immuno-properties for potential Tuberculosis diagnosis and vaccine candidates

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Background

Mycobacterium tuberculosis (Mtb) is a causative agent for Tuberculosis (TB) with estimated a quarter of the world population has been infected and has the highest cumulative mortality rate among other infectious diseases with 1.3 million death in 2020 alone (WHO, 2021). The report also highlighted TB as the leading cause of mortality by single infectious agent. Ten percent of Latent TB Infection (LTBI) will develop into active infection. Existing diagnostic methods e.g. Tuberculin Skin Test and are not accurate. Newer technique i.e. Interferon Gamma Release Assay is expensive, thus hardly affordable especially in low to middle income countries with high TB incident rate. Currently, there are lack of cheap, sensitive, and specific diagnostic methods to diagnose individuals with LTBI and the current BCG vaccine has limited coverage against adult TB infection.

Methodology

Reversed vaccinology technique was applied to shortlist potential *Mtb* T-cell epitopes (TCEs) for diagnosis and vaccine candidates. Six immuno-criteria were selected to rank the TCEs i.e. matching to *Mtb* H37Rv antigens, coverage across 273 *Mtb* strains, association with experiment types of highly expressed *Mtb* genes, promiscuous epitopes in terms of loci and alleles, and population coverage. Each criterion dataset was obtained from a different database and individually preprocessed, analysed using a selected prediction method, and collated using an in-house script. Each factor was equally weighted, and all of the factor results were integrated into a relational table to be used as an interactive scoring platform. At the end, each TCE has six scores from the six factors, and they are combined to produce the final score.

Results and Discussion

Looking from the perspective of individual criterion: 1) almost all TCEs have 1-to-1 matching to Mtb H37Rv antigens; 2) They also have a high matching coverage across all 273 completed strains tested; 3) Majority of them matched to gene used in human-related experiment literature; 4) Almost all of the TCEs bind to at least a locus, in average a TCE bind to 3.4 (out of 8) loci, and restriction to A-B-C loci has the highest TCE count; 5) In average, each TCE is restricted to 26 (out of 143) alleles; 6) They were tested with their coverage in world, Asian, Southeast Asia and Malaysia populations; where more than half of TCEs cover more than 90% of the world population. Looking from the perspective of top scoring TCEs, top thirty TCEs were further analysed for their associations to cellular functional groups, sequence motifs, common genes used for diagnosis and vaccine and Mtb Bcell epitopes. Most of the shortlisted epitopes were found to be parts of potential genes (encoded for proteins) that are widely studied for their cellular and humeral body response i.e. ESAT-6 family (e.g. esxJ, esxK, esxM, esxP, esxW, espJ, and espK), Ag85B, Ag85C, and HSP16.3.

Conclusion

These genes have been showed to have potential in TB diagnosis (e.g. Interferon Gamma Release Assay technique) and vaccine candidates.

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0-31

Desiccation tolerance mediates ST239-SCCmec type III-SCCmercury to ST22-SCCmec type IV MRSA clonal replacement in hospital settings

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ST239-SCCmec type III-SCCmercury (ST239-III) to ST22-SCCmec type IV (ST22-IV) clonal replacement in methicillin-resistant *Staphylococcus aureus* (MRSA) has been reported globally in many hospitals during the first decade of the new millennium, including in Hospital Canselor Tuanku Muhriz (HCTM), Kuala Lumpur, Malaysia. It was hypothesized that the clonal replacement occurred due to faster growth rate of ST22-IV [1]. To confirm this, we studied and compared the competitive fitness of representative strains from the two clones, ST239-III and ST22-IV isolated from HCTM via broth co-culture and desiccation experiments.

Methodology

The strains used in this study were M57/2005 and M222/2009 (both ST239-III), M181/2017 and M080/2017 (both ST22-IV). Chloramphenicol (20ug/ml) susceptibility was used to differentiate the STs of tested strains; ST239 strains were resistant, ST22-IV strains were susceptible. In broth co-culture experiments, pairs of tested strains from different STs were cultured together for 48 hours in brain heart infusion (BHI) broth and platted on drug-free and chloramphenicol-supplemented BHI agar. In desiccation experiments, pairs of tested strains from different STs were exposed to desiccation for 48 hours on petri plates prior rehydration of the cultures with normal saline, followed by drug-free and chloramphenicol-supplemented BHI agar platting and incubation at 37°C overnight. Numbers of colonies on agar plates for both STs in both experiments were enumerated. SCC*mec* typing of representative colonies was also performed for ST confirmation.

Results and Discussion

For broth co-culture experiments, mean of 194.5, (135-272) ST239-III colonies and 31.5, (4-58) ST22-IV colonies were observed at 10⁻⁶ serial dilution. On the other hand, ST22-IV produced a higher number of surviving colonies on desiccation assays (29, 5-58) compared to ST239-III (2.5, 0-5) at 10⁻⁶ serial dilution. While ST239-III outcompeted ST22-IV in all tested pairs for the broth co-culture studies, the latter clone had better tolerance to desiccation. As MRSA is a nosocomial pathogen with clonal dissemination in hospitals [2], higher desiccation tolerance could have provided ST22-IV an edge to survive and disseminate, ultimately replacing ST239-III in the hospital environment.

Conclusion

ST22-IV was found to have higher desiccation tolerance compared to ST239-III. This edge in bacterial fitness might be the cause of ST239-III to ST22-IV clonal replacement in nosocomial MRSAs; further studies will be required to confirm this.

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O-33

Ligand-based pharmacophore modeling, molecular docking and molecular dynamics study targeting prolyl oligopeptidase enzyme for effective treatment for Parkinson's disease: Computational approach Yahaya Sani Naiib^{1, 2}. Yusuf Oloruntovin Avipo^{1, 3}. Waleed Abdullah

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From The International Conference on Molecular Diagnostics & Biomarker Discovery 2022 (MDBD 2022) Penang, Malaysia. 11 – 13 October 2022 Prolyl oligopeptidase (POP) are serine protease enzymes implicated in the pathogenesis of Parkinson's disease (PD) through the increased aggregation of α-synuclein protein in the brain. The current treatment options for PD are only symptom-targeted, while an effective therapeutic strategy remains a challenge. The study aimed to identify potent anti-PD drugs with inhibitory potentials against POP using ligand-based receptor modelling, Glide XP docking, molecular dynamics and pk-CSM pharmacokinetics ADMET prediction parameters. Results indicated that the ligand-based (LB) model generated pharmacophore with 6 features, having 1 hydrophobic, 1 positive ionizable group, 2 aromatic rings, and 2 hydrogen bond acceptors. A total of 23 hits with a Gunner-Henry (GH) score of 0.7 and enrichment factor (EF) of 30.24 were obtained as validation protocols, making it an ideal model. The LB model retrieved 177 hit compounds from 69,543 natural Interbioscreen databases virtually. Interestingly, ligands 1, 2, 3, 4 and 5 demonstrated higher binding affinity to the enzyme with Glide XP docking of -9.004, -8.818, -8.749, -8.708, -8.707 kcal/mol than GSK552 and ZPP with -8.187, and -6.835 respectively. Similarly, their binding free energies were -48.424, -51.879, -51.684, -45.191, and -49.567 kcal/mol respectively. Molecular dynamics indicated that the ligands 1, 2, & 4 demonstrated better stability than GSK552. Pharmacokinetic profiles of the ligands indicated their druggability and low toxicity profile. The ligands are recommended as adjuvant /single candidates as ant-PD candidates for translational study.

0-34

Inhibitory effects of andrographolide in PC-3 cell line and the induction of apoptosis via the involvement of caspases activity

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Background

Andrographolide is a labdane diterpenoid isolated from the plant *Andrographis paniculata*. This substance has numerous medicinal uses, notably anticancer effects. A previous study has revealed that andrographolide inhibits the growth of lung, brain, colon, and breast cancer cells [1]. Due to a lack of research, however, it is thought that the knowledge of andrographolide's anti-cancer effects on prostate cancer cells is relatively poor. In the current study, andrographolide was assessed on PC-3 cells, an aggressive androgen-independent prostate cancer cell line.

Methodology

Cytotoxicity analysis is vital in drug discovery research for assessing the biocompatibility of the drug being used on cancer cells. This work used the WST-1 assay to determine the cell survival of PC-3 cancer cells and Hs27 normal cells exposed to varying doses of andrographolide (0-200 μM). Metastasis is essential for the disease to progress; hence, the scratch assay and the transwell invasion assay were used to test andrographolide on PC-3 cells. In order to study the cell death mechanism after the PC-3 cells are treated with andrographolide, DNA damage and apoptosis assay have been done. Finally, the caspase-mediated pathway is analysed by carrying out caspase 3, 8 and 9 assays upon the treatment of andrographolide.

Results and Discussion

The results indicate that andrographolide dose-dependently suppresses the viability of PC-3 cells but not Hs27 cells. The NCI considers the LC 50 value of 26.42 μM (after 48 hours of incubation) acceptable. In this study, three different concentrations of andrographolide were used: control, half LC 50, and LC 50 (0, 13.21, and 26.42 $\mu\text{M})$ in all subsequent analyses. The results showed that andrographolide inhibits both migration and invasion compared to the control. The presence of a comet tail has revealed that a 26.42 μM andrographolide treatment produces the maximum DNA damage to single cells, followed by 13.21 μM and 0 M. In addition, the maximum cell death activity was seen at 26.42 M andrographolide concentration, followed by 13.21 μM and the control. The activity of caspases 3

(executor caspase), 8 (intrinsic pathway), and 9 (extrinsic pathway) in mediating apoptosis increased significantly. The half LC 50 (13.21 $\mu\text{M})$ demonstrates significantly more activity in these caspases than the LC 50 (26.21 $\mu\text{M})$, in accordance with the new finding related to the caspase storm scenario [2]. This enables us to identify 13.21 μM as the andrographolide dosage that approaches the ideal range for caspase activity activation.

Conclusion

The ability of andrographolide to prevent the progression of cancer in PC-3 cancer cells has been proven through the regulation of caspase-mediated apoptosis, suppression of metastasis, and induction of DNA damage.

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0-35

Lentiviral modification of hard-to-transduce NK-92MI cells

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Background

Natural killer (NK) cells are lymphocytes that play an important role in both our innate and humoral immunities. NK cell functions are dictated by the myriad of receptors, regulating both activating and inhibitory signals. Studies found that through interactions with the CD16 receptor alone, NK cells can be activated for antibodydependent cellular cytotoxicity (ADCC) to occur. The ADCC mechanism can be used to home NK cell killing towards designated targets through detection by target-specific antibodies. Unfortunately, NK cell studies are hampered by the tedious sourcing of primary NK cells. NK cell lines such as NK-92 and NK-92MI are now available but they lack certain features compared to primary cells. Furthermore, NK cells are less susceptible to genetic modifications such as electroporation and viral-based methods. Recently, lentiviral vectors were successfully used in the clinical setting for cell modifications. In this study, we used a third-generation lentiviral vector, termed RRLlentivirus, to modify the NK-92MI cells for surface expression of CD16 receptors and possibly restore their ability for ADCC.

Methodology

High binding affinity 158V-CD16 and GFP gene sequences were molecularly cloned into RRL-lentiviral transfer vector. The transfer vector plasmid containing the gene of interest and three types of helper plasmids were simultaneously transfected into HEK-293T packaging cells by calcium phosphate transfection. After 48 hours, the supernatant containing the lentiviral particles was collected and filtered through a 0.45um CA filter. This was then concentrated using PEG-6000 precipitation method and stored in serum-free media at -80°C. Using K-562 cells as positive controls, NK-92MI cells were transduced using GFP-RRL lentivirus or CD16-RRL lentivirus. Transduction replicates were done with or without the addition of polybrene polycation. The expression of reporter GFP was monitored using fluorescence microscopy, and the levels of transgene expressions were further validated through flow cytometry using FITC-conjugated anti-human CD16 antibodies.

Results and Discussion

After GFP-lentivirus transduction was performed, the control K-562 samples showed GFP fluorescence on day 3, while the NK-92Ml samples displayed GFP on day 5. The difference between the samples

transduced with and without polybrene was compared through flow cytometry. At MOI of 1, GFP-transduced K-562 cells resulted in a GFP-positive population of 95.6% without polybrene and this improved to 99.5% when polybrene was added. At the same MOI, CD16-transduced K-562 did not result in the formation of two distinct CD16 populations. However, all transduced samples showed a significant positive shift in the detected FITC compared to negative controls. At the MOI of 10, GFP-transduced NK-92MI cells resulted in GFP-positive population of 3.5% without polybrene, and the population increased to 7.2% with polybrene added. CD16-transduced NK-92MI cells resulted in a CD16-positive population at 1.0%, while the addition of polybrene increased this expression to 3.1%.

Conclusion

This study demonstrated the feasibility of using a third generation RRL-lentiviral vector for NK-92MI cell transduction, but further optimisations will be required before *in vitro* assays such as natural cytotoxicity and ADCC assays can be performed using these modified cells.

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0-37

Immunoinformatics analysis on human coronavirus spike protein for universal immunogen discovery

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Background

Coronaviruses are well-known to possess high mutation rate. The recent pandemic has demonstrated this fact that multiple SARS-CoV-2 variants have emerged since the first occurrence in 2019. Variants such as Beta, Delta and Omicron variant, even showed the ability of immune evasion in convalescent and vaccinated individuals [1], raising global concern about the efficacies of existing vaccines. Immunoinformatics approach is gaining traction in vaccine development due to significant time and cost reduction in immunogenicity studies and improved reliability [2]. Viral genome can be analysed for the mapping of potential T-cell and B-cell epitopes. Structural proteins, particularly spike protein (S) have been studied extensively as promising vaccine candidates. One rationale is that the RBD of SARS-CoV-2 attaches to the ACE2 receptor on the host cells to initiate infection. Taken together, a vaccine that offers protection over a wide spectrum of coronaviruses is crucially demanded.

Methodology

The sequences of S proteins of the SARS-CoV-2 variants as well as SARS-CoV and MERS-CoV were retrieved from the NCBI database. Multiple sequence alignment (MSA) was performed to identify the conserved regions among S proteins from different human coronaviruses. All conserved regions were analysed for the antigenicity through VaxiJen. The conserved regions that passed the threshold value of 0.5 were then analysed for T-cell and B-cell epitope

predictions using NetMHCpan EL4.1, IEDB recommended 2.22 and BepiPred 2.0, respectively. The antigenic conserved regions were also used to construct 3D model using SWISS-MODEL followed by structural refinement using GalaxyRefine2. All refined models were validated using ERRAT and PROCHECK. Lastly, the docking of 3D models against TLR-3 was performed via PatchDock and FireDock.

Results and Discussion

Based on the MSA result, S protein contains 12 conserved regions, which were then subjected to further analyses. Altogether 9 conserved regions were above the antigenicity threshold value and therefore selected. In terms of epitope prediction, the antigenic conserved regions were predicted to contain a total of 69 MHC Class-I epitopes, 45 MHC Class-II epitopes and 5 linear B-cell epitopes. Furthermore, all antigenic conserved regions were sent for 3D model building followed by structural refinement. The best refined model for each conserved region was chosen based on the Galaxy energy. These refined models were validated and the results showed that the models were of good quality. Following that, the refined model for each conserved region was docked against TLR-3. The global energies of complexes formed between the conserved regions and TLR-3 fell within the range of -11.74 to -7.36 kcal/mol. This implies that the conserved regions have good binding affinities with TLR-3.

Conclusion

The identified conserved regions of S protein were predicted to have a significant number of epitopes and showed promising docking results. This study provides some insights about the interaction of conserved S peptides with TLR-3, contributing to the vaccine design. Still, further analyses such as molecular dynamics and immune simulation are required to polish the results. In vitro and in vivo validation are also essential to evaluate the immunological roles of designed universal coronavirus vaccine.

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0-38

Andrographolide induced apoptosis by enhancing c-Myc/p53 in human glioblastoma DBTRG-05MG cell line

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Background

Human glioblastoma multiforme (GBM) is one of the most malignant brain tumors. The conventional GBM treatment faces a problem due to the presence of the blood-brain barrier (BBB) and the post-treatment effect. Then, andrographolide has attracted many researchers because it can cross the BBB and easily distribute it into different brain regions [1]. Next, targeted therapy also is one of the treatments that can reduce the post-treatment effect because it can specifically target cancer cells without affecting normal cells. Recently, the small molecule targeted therapy by exploring on the molecular targets of oncogenes or tumor suppressor genes such as c-Myc and p53 in regulating the apoptosis signaling pathway still remains debatable [2]. Therefore, this study aimed to elucidate the mechanism action of andrographolide towards c-Myc/p53 induced apoptosis in DBTRG-05MG cell line.

Methodology

The cytotoxic effect of andrographolide was assessed by WST-1 assay in DBTRG-05MG and SVGp12 cell lines. Both cell lines were treated for 24, 48, and 72 hours with varying concentrations of andrographolide (0.781 – 200 μM). The WST-1 reagent was added to each well. Then, plates were incubated, and the absorbance (Abs.) was measured with an ELISA reader (Multiscan Spectrum) at wavelength 450 nm and the reference reading of 630 nm. The LC₅₀ values were computed and will be further used as active doses and time in the subsequent experiments. Next, the apoptosis assay was assessed using annexin V-FITC / PI double staining. Cells pellet of DBTRG-05MG were collected after treatment with andrographolide and resuspended with V-FITC, and PI staining. The cells were incubated in the dark at room temperature for 30 minutes before analyzing the stained cells using a BD FACS CantoTM Flow Cytometer. The gene and protein expression level c-Myc and p53 signaling pathway were then assessed using qRT-PCR and western blot. The protein-protein interaction between c-Myc and p53 was determined by a reciprocal experiment of the co-immunoprecipitation (co-IP) using DBTRG-05MG total cell lysate.

Results and Discussion

Andrographolide has significantly reduced the viability of DBTRG-05MG cell lines in a concentration- and time-dependent manner. The recorded LC_{50} values for DBTRG-05MG cell lines for 24, 48 and 72 hours of treatment were 42.82 μ M, 27.21 μ M and 13.93 μ M, respectively. While in the non-cancerous cell line, SVGp12 cannot be determined because the percentage cell viability for the normal cell remained ~80% after being treated with the highest concentration (200 µM) of andrographolide for 24h results. Andrographolide also induced apoptosis in the DBTRG-05MG cell line, by inducing c-Myc and p53 expression at the gene and protein level. The percentage of apoptotic cells in control cells was 0.01%, and after the cell lines were exposed to 13.95 µM and 27.9 µM of andrographolide for 72 hours, the percentage of the apoptotic cell increased to 5.2% and 16.5%, respectively. Western blot results demonstrated that c-Myc overexpression also increased the production of the anti-apoptotic protein p53. Our findings revealed that c-Myc and p53 positively interact in triggering the apoptotic signaling pathway when the lane immunoprecipitated (IP) result for p53 and probed with anti-c-Myc in western blot analysis, which showed the presence of the band for c-Myc protein (62kDA).

Conclusion

This study successfully discovered the involvement of c-Myc and p53 in the inhibition of DBTRG-05MG cell line via apoptosis following andrographolide treatment. However, further study needs to be extended to evaluate the cytotoxicity of andrographolide in vivo and in vitro. The function of other regulatory genes that may interact with the c-Myc signaling pathway must also be investigated intensively.

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0-39

In vitro uptake and activation of human dendritic cells by liposomes derived from total lipid of Mycobacterium smegmatis

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Liposomes are self-assembled lipids that have attracted scientific attention as efficient drug delivery vehicles and adjuvants. They possess a unique vesicular structure and can be derived from natural and synthetic substances. Liposomes can mimic the biological membrane of drugs, thus extending the half-life and minimizing the toxicity levels while delivering them to the target organs. The adjuvant mechanism of liposomes has been intimately associated with the stimulation of desired immune responses upon the exposure of antigen and immune cell targeting. This current study mainly targets to investigate the uptake and activation of human dendritic cells (DCs) by liposomes derived from the total lipid of *Mycobacterium smegmatis*.

Methodology

Liposomes derived from *Mycobacterium smegmatis* were synthesized and characterized using Ziehl-Neelsen (ZN stain) and field emission scanning electron microscopy (FESEM). Fresh whole blood has been collected from three distinctive groups: TST-negative individuals, TST-positive individuals, and active pulmonary TB patients. Peripheral blood mononuclear cells (PBMCs) were separated by Lymphoprep density centrifugation and cultured with lipopolysaccharide (LPS) and liposomes. The activation of DCs by *M. smegmatis* liposomes has been analyzed through the expression level of DCs surface markers (HLA-DR, CD11c, CD123, CD86) in flow cytometry and the secretion of cytokines (IL-4, IL12p70, and IFN-γ) via ELISA.

Results and Discussion

The characterization of liposomes under FESEM demonstrated a size ranging from 20 nm⁻¹³⁵ nm with spherical structures. The results showed levels of HLA-DR and CD86 expression reduced in the group of DCs of active pulmonary TB with liposomes compared to the group of TST-negative and TST-positive individuals. However, the interaction of DCs from active pulmonary TB patients with liposomes exhibited an increased level of HLA-DR with low expression of CD86 compared to other groups. IL-12p70 was a highly significant increase in active pulmonary TB group compared to the group of TST-negative and TST-positive individuals. The group of active pulmonary TB patients also showed a significant increase in IL-4 and IFN-γ than the group of TST-positive individuals. Meanwhile, the secretion of all cytokines was the lowest in the group of TST-positive individuals compared to the other groups. These events emphasized the malfunction of DCs in a TB condition and highlighted the capability of liposomes to support the antigen presentation by human DCs in TB infection.

Conclusion

Liposomes synthesized from *Mycobacterium smegmatis* had been successfully produced and classified as small unilamellar vesicles (SUVs). The exposure of DCs with liposomes particularly improved the antigen presentation activity with low activation of DCs in active pulmonary TB patients compared to the other study cohort. The results from this current study supported liposomes as potent vaccines and adjuvants for immunotherapy.

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0-40

Identification of medicinal fungi by molecular analysis

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Background

Mushrooms or higher fungi are heterotrophic eukaryotic organisms on Earth. Fungi are also having its great medical and environmental importance. Hence, the precise and rapid identification of fungi species

from specimens is important for the field applications, such as, sustainable use and conservation of biodiversity, prevention and control of fungal pathogens, human health and ecological monitoring [1]. DNA barcoding is a great tool for specimen identification that can be used for identifying and recognizing various types of fungi. The internal transcribed spacer (ITS) regions of the nuclear ribosomal RNA gene cluster were utilized as a universal DNA barcode marker for fungi specimens which was suggested by Schoch and his team [2]. In this study, we identified 11 randomized fungi specimens by using ITS primers.

The fungi specimen grown on Petri plates containing PDA medium was obtained from the lab. A small number of mycelia of fungi specimens was picked using sterile forceps from the fully-grown culture and was transferred into a 1.5 mL microcentrifuge tube consisting of 600 µL of sterile distilled water. The mycelia were homogenized by using pipette tips. The mixture was vortexed thoroughly and centrifuged at 10,000 x g for 1 minute. The supernatant was discarded and continued by using the EasyPure Plant Genomic DNA kit according to the manufacturer's instructions to extract fungi DNA. MyTaq HS Red Mix, 2x, universal ITS primer pairs, extracted fungi DNA sample template and sterile distilled water were used for performing PCR amplification using Agilent SureCycler 8800 with 35 cycles of amplification. The PCR products were undergone electrophoresis to separate DNA. The gel was observed and captured using a Syngene Bioimaging system. The EasyPure PCR purification kit was used to purify PCR products to remove primers and nucleotides according to the manufacturer's instructions. The purified PCR products were sent for sequencing. The FinchTV software, BLASTn algorithm at the NCBI website and DNA Bold program were used to analyze the sequences data.

Results and Discussion

Eleven fungi specimens were determined by its size by using the Syngene Bioimaging System. The highest base pair of fungi specimens was Sample 10, which had 850 base pairs. The lowest base pair of fungi samples were Samples 1, 4, 5 & 6, which had 600 base pairs. The range of base pair of fungi specimens was from 600 bp to 850 bp. Therefore, 11 fungi specimens were identified by using the FinchTV software, BLASTn algorithm at the NCBI website and DNA Bold program through analyzing their sequences data. The outcomes of the analysis revealed these specimens were *Hypoxylon sp. LA01, Lignosus rhinocerotis, Ganoderma lucidum, Cordyceps militaris, Hemistropharia albocrenulata, Inonotus obliquus,* and *Hericum erinaceus*.

Conclusion

The DNA extracted from randomized fungi specimens were identified successfully using molecular analysis. The works presented here reveal this method will potentially aid in the authentication of medicinal fungi from unknown specimens.

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0-41

Effect of andrographolide on proliferation, migration, and invasion of MDA-MB-231 and MCF-7 cell lines

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Breast cancer is the commonly diagnosed and one of the leading causes of death among women worldwide [1]. Current standard treatment includes chemotherapy, surgery, and radiotherapy. Some patient discontinued their treatment due to the side effects of the treatments which leads to relapse. Therefore, alternative medicines from natural plants with fewer side effects now widely studied. Andrographolide is a diterpene lactone derived from Andrographis Paniculata. Andrographolide had previously demonstrated to have anti-inflammatory, anticancer, anti-fungal, and anti-viral properties. This study is done to determine anti-cancer effects of andrographolide in breast cancer cells by evaluating the cell viability, migration, and invasion in two different cancer cell lines which are MDA-MB-231 (a triple negative cell line) and MCF-7 (ER positive cell line).

Methodology

MDA-MB-231 and MCF-7 cells were treated with a different concentration of andrographolide (0.781 μ M - 200 μ M) for 24, 48 and 72 hours. The cell viability was observed using the WST-1 reagent. The LC $_{50}$ and ½ LC $_{50}$ values obtained from WST-1 were then used for scratch and transwell invasion assays to determine the rate of migration and invasion for both cell lines. The cells were both treated with LC $_{50}$ and ½ LC $_{50}$ concentration of andrographolide for a different timeline, MDA-MB-231 was treated for 72H and MCF-7 was treated for 48H before proceeding to scratch and invasion assay.

Results and Discussion

Andrographolide had shown a significant effect on MDA-MB-231 and MCF-7 cells by suppressing the cell viability in a concentration dependent manner. Our findings are consistent with the previous study that showed andrographolide reduced the cell viability in cancer cell but have a minimum effect on normal cells in a concentration dependent manner. Andrographolide inhibit the migration of MDA-MB-231 and MCF-7 cell lines at 72 and 48 hours, respectively. The wound gap area increased in both cells treated with andrographolide at two different concentrations compared to both untreated cells. Invasion assay showed that andrographolide inhibit the invasion of MDA-MB-231 and MCF-7 cell lines at a low concentration which aligned with the previous research that showed andrographolide inhibit the invasion of MDA-MB-231 [2].

Conclusion

Andrographolide showed the migration and invasion inhibition properties in a concentration dependent manner in MDA-MB-231 and MCF-7 cells. Therefore, andrographolide is shown to have a promising future to be develop as a potential treatment for breast cancer. Further research needs to be done to elucidate the mechanism of cell death and whether it involved in either cell cycle or apoptosis pathway.

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0-42

Culture and biochemical testing versus 16S rRNA next-generation sequencing for bacterial identification from clinical samples: Practicability, cost and turn-around time in a Malaysian laboratory Nurnabila Syafiqah Muhamad Rizal¹, Hui-min Neoh¹, Ramliza Ramli¹, Petrick @ Ramesh A/L K Periyasamy¹, Alfizah Hanafiah¹, Muttaqillah Najihan Abdul Samat¹, Toh Leong Tan¹, Kon Ken Wong¹, Sheila Nathan¹, Sylvia Chieng¹, Seow Hoon Saw² and Bee Yin Khor³

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Background

Bacterial pathogens are mostly identified via culture and biochemical testing (CBtest) in Malaysian clinical microbiology laboratories. In this study, we assessed the practicability, cost and turn-around-time (TAT) of utilising 16S rRNA next- generation sequencing (16SNGS) versus CBtest for bacteria identification from clinical samples collected at the Department of Diagnostic Laboratory Services, UKM Medical Centre, Kuala Lumpur.

Methodology

Twenty-four clinical samples (eight each from pus, respiratory and urine specimens) were included into the study. CBtest was carried out via microscopy and staining the samples, bacterial culture and isolation followed by biochemical tests for species identification. 16SNGS was performed via amplification of V3- V4 regions of 16S rRNA gene from total DNA extracted from each sample and sequenced using the MiSeq system; raw sequencing data was analyzed using the B.E Patho software. TAT was calculated from the initiation of sample processing to test report generation. Cost analysis included cost for reagents and disposable procurement as well as handson labour charges.

Results and Discussion

Via CBtest, 18 samples were reported as positive for single- bacterial infection (n = 5, 20.8%), mixed growth (n = 8, 33.3%) and normal flora (n = 5, 20.8%), respectively; remaining samples were culture-negative. 16SNGS detected the relative abundance of various bacteria genera in all tested samples. TAT of 16SNGS workflow (six days) was shorter compared to CBtest for identification of slow-growing, fastidious bacteria (72 days). Utilisation of the B.E. Patho "plug-and-play" bioinformatics suite allowed 16SNGS analysis to be performed by personnel without bioinformatics training. Nonetheless, identification cost per sample via 16SNGS (RM 1704.70) was very much higher than CBtest (RM 2.80 – RM 42.90) for all samples.

Conclusion

16SNGS allowed identification of various bacterial genera in clinical samples with a faster TAT for slow-growing and fastidious bacteria. Lowering of sequencing costs will facilitate future adoption of the platform in clinical microbiology laboratories.

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0-43

Isolation and production of recombinant monoclonal antibody proteins against a *Toxocara canis* antigen using phage display technology Zamrina Baharudeen and Anizah Rahumatullah

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Toxocariasis is a neglected zoonotic parasitic disease which mainly caused by *Toxocara canis*, an intestinal parasitic roundworm of dog. Human toxocariasis has a global distribution, with people from socioeconomically deprived populations more adversely affected. The limitations of diagnosing human toxocariasis is the signs and symptoms of the disease are non-specific and similar to other helminth infection [1]. This may lead to misdiagnosis thus underestimating the actual global impact of the disease. Existing serology-based detection method mainly rely on IgG antibody-based assays that suffers from the difficulty in differentiating past and active infections. The assays also lack of high diagnostic specificity due to cross-reactivity with antibodies to other helminths [2]. In the present study, novel recombinant monoclonal antibodies were isolated and the binding of the proteins were verified.

Methodology

Monoclonal antibody against recombinant *Toxocara* excretory-secretory 26 antigen (rTES-26) was isolated utilizing previously reported helminth phage display library via biopanning. Polyclonal and monoclonal phage ELISA were carried out at the end of selection. Positive clones were sent for sequencing and the scFv antibody clones verified using IMGT/V-QUEST bioinformatics tool. Selected positive monoclonal antibody clones with complete scFv gene were subcloned into pET-51b(+) and transformed into SHuffle® T7 Express *Escherichia coli* host cell. Antigen-antibody binding assays were performed using rTES-26 and native *T. canis* antigens.

Results and Discussion

The significant increase in absorbance reading of polyclonal ELISA (OD₄₀₅ ranging from 0.321 to 3.020) indicating rTES-26 specific phage antibodies successfully enriched. In total 384 individual clones were screened during monoclonal ELISA and five positive binders were identified with absorbance values ranging from 0.45 to 3.91. Three clones from distinct gene families (Ab 48: IgHV3-LV1; Ab 49:IgHV3-LV3; Ab 50: IgHV6-LV3) were selected, however only two clones: Ab 48 and Ab 49 demonstrated complete insertion of the full-length scFv antibody sequence following sub-cloning. Both clones were successfully expressed and purified at satisfactory levels in terms of purity and yield. The antigen-antibody binding analyses showed that both Ab 48 and Ab 49 able to bind the recombinant and native form of Toxocara protein. However, in total Ab 49 showed better performance in protein yield and binding affinity compared to Ab 48. Hence, the enriched monoclonal antibodies from the immune libraries shows to have varying binding affinities and specificities [3].

Conclusion

In summary, both antibody clones showed their diagnostic potential and the binding evaluation suggest that Ab 49 has the potential to be used as an efficient tool for human toxocariasis diagnosis.

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0-4/

Optimization of transient expression of recombinant IgG binding protein, $Fc\gamma RIIa$

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Background

IgG Fc binding proteins (FcγRs) are immunoglobulin receptors that particularly bind to the human IgG (hlgG) molecules through its Fc portion. Among the members of FcγRs, FcγRllA is slowly gaining interest from scientific community due to its different isotype specificities and the significant binding affinity towards IgG2. Current technology has presented the production of recombinant proteins in mammalian cell lines to introduce appropriate protein folding and post-translational modifications [1]. However, the production of recombinant proteins using mammalian expression systems is laborious, time-consuming, and costly [1]. Therefore, this study provided a simple protocol for optimized transient transfection and expression of recombinant IgG binding proteins to achieve a high yield in a short span of time.

Methodology

FreeStyle™ 293-F cells in the prewarmed FreeStyle™ 293 Expression Medium were incubated in a 37°C incubator containing a humidified atmosphere of 5% CO₂ in air on an orbital shaker platform rotating at 120 rpm. On the day of transfection, the cells were suspended in a volume of prewarmed medium to obtain a concentration of 1.0×10⁶ cells/mL. For each mL of transfection volume, $1\mu g$ of plasmid DNA and $2\mu L$ of 293fectin diluted in Opti-MEM were needed. After adding the DNA/293fectin mixture, the suspension cells were incubated for 96 hours. The cell pellets were harvested by spinning down the cells for 5 mins at 300 xg and the intracellular protein was harvested by using lysis buffer. After the protein has been purified using immobilized metal affinity chromatography (IMAC), the protein was subjected to sodium dodecyl sulphatepolyacrylamide gel electrophoresis (SDS-PAGE) and Western Blot analyses for the visual evaluation of expression. The protein concentration was measured by using Nanodrop Spectrophotometer.produced in less than one week. In general, the recombinant proteins with expression levels up to 100 mg/L can be obtained. Since the expression level may differ depending on the nature of the recombinant proteins [2], the transfection method had been refined by varying the incubation time posttransfection, ending up with an ideal harvest time of 96 hours. The harvested protein was purified by using IMAC for downstream analysis. In SDS-PAGE analysis, an intense protein band at approximately 38 kDa had been observed, indicating the expression of the recombinant proteins. Meanwhile, in Western blot analysis, anti-histidine antibody was used to detect the hexa-histidine tagged recombinant protein.

Conclusion

An optimized protocol that serves as a general guide for the transient transfection and expression of recombinant IgG binding protein, FcyRIIA had been described. The simplicity of this protocol had exhibited an efficient transfection and a fast yet high-yield expression of proteins.

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0-45

Cloning of IgM Fc receptor for mammalian expression system

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Mammalian expression system has become routine in the production of pharmaceutical protein as it offers post-translational modifications (PTMs) which allows production of complex proteins and glycosylated enzymes where their native structures and activities are preserved. In order to express the recombinant protein in mammalian cell lines efficiently, selection of an appropriate expression vector that carry the interested recombinant gene plays an important role. Polymeric immunoglobulin receptor (plgR), an lgM Fc receptor is chosen to be expressed due to its various clinical potentials such as the possibility to serve as predictive biomarker target owing to its correlation with various cancer malignancies, potential role, and underlying mechanisms in body immunity [1, 2].

Methodology

The plasmids that carried *PIGR* gene were purified from the bacterial overnight culture, followed by polymerase chain reaction (PCR) to amplify the interested DNA segment. The PCR products and mammalian expression vector (pcDNA5/FRT) were then subjected to restriction digestion and ligation process prior to bacteria transformation. The final product was sent for sequencing before the mammalian cells transfection.

Results and Discussion

The presence of bacterial colonies in ampicillin-containing agar plate indicating the success of bacteria transformation. The cloning of the gene of interest into the mammalian expression vector (pcDNA5/FRT) was then verified through restriction digestion, showing two bands where both represented the vector backbone (5002bp) and PIGR gene sequence (2490bp) respectively. The sequence was checked for any point mutation through comparison of each nucleotide with the reference sequence. Flp-In System involves the introduction of Flp Recombination Target (FRT) site into genome of both expression vector and mammalian cell line of choice. The expression vector that carrying the gene of interest will integrate into the host cell genome through Flp recombinase-mediated DNA recombination at the FRT site. This system allows stable expression cell lines to be created, providing advantages such as rapid and efficient protein expression by the subsequent generation of Flp-In cell lines [3].

Conclusion

Gene cloning is an important step for the protein expression by living cells, especially for mammalian cells. The quality and sequences of the recombinant plasmid will impact the efficiency of protein expression. In future, the constructed recombinant plasmid (pcDNA5/FRT-PIGR) will be transfected into chosen mammalian cell line to express the recombinant protein as the sequence were checked accordingly. Antibiotic selection will be performed to select cells where the plasmids are integrated into genome successfully. Single cells that are able to survive upon transfection and selective growth will be expanded to generate a clonal population. Stable clones are selected after screening and ready for protein production.

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0-46

New β -carboline compound as a promising anticancer agent in chronic myelogenous leukemia (CML)

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Background

Cancer is known as one of the serious causes of deaths among the world population and it is believed that the number of cases to rise over the years. Chemotherapy drugs are considered as the most reliable and common treatment for cancer. However, these drugs have limitations in terms of effectiveness and severe side effects. In addition, drug resistance is another factor that hinders successful treatment rate among patients. Therefore, finding a lead compound for cancer which addresses these issues are of utmost important. In the constant search for new anticancer agent alternatives, β -carboline has been identified as a good candidate due to its various reported health indications, particularly in anticancer. Therefore, the aims of the study are to investigate the potential of a newly synthesized β -carboline compound in eliciting anticancer activity in chronic myelogenous leukemia (CML) and to determine the apoptosis mechanism induced by the compound.

Methodology

β-carboline compound (M25) was synthesized and confirmed using gas chromatography-mass spectrometry (GC-MS) analysis. MTT assay was conducted to evaluate cytotoxicity effect of M25 against CML cells (K562 cell line). Toxicity of M25 was determined by calculating its selectivity index (SI), through human fibroblasts (Hs27 cell line) and mouse fibroblasts (BALB/c3T3). Mode of cell death induced by M25 was further evaluated through the apoptosis assay using flow cytometry analysis with Annexin V-FITC and Propidium iodide (PI). Lastly, *in silico* analysis was conducted to study the binding affinity between M25 and identified proteins involved in the apoptosis pathway using Discovery Studio software.

Results and Discussion

β-carboline compound M25 was synthesised with high purity and high yield. M25 induced cytotoxicity in K562 cell line with a low IC₅₀ value of 0.8 μM. This indicated high potency of M25 in killing human CML cells. Based on IC₅₀ values for non-cancer cells Hs27 and BALB/c3T3, the calculated SI were 16 and 25 respectively. The high selectivity showed that M25 is highly specific in killing CML cells, thus reflecting good safety profile for the compound. Flow cytometric analysis also showed that significant number cell death induced by M25 compound was due to apoptosis. In understanding proteins involved in the apoptosis pathway, molecular docking study investigated the binding affinity of M25 as ligand with apoptosis-related proteins such as apoptosis inducing factor (AIF), endonucleases G (Endo G), caspase 3 (CAS 3), caspase 9 (CAS 9) and caspase 8 (CAS 8). Interestingly, the highest binding affinity was recorded for AIF and Endo G, which suggested a unique caspase-independent pathway of apoptosis.

Conclusion

The newly synthesized M25 has excellent potential as an anti-leukemic agent as it has shown high cytotoxicity activity and selectivity for K562, thus indicating good effectiveness and safety profile. In addition, in silico study suggested that M25 induced caspase-independent

apoptosis, with the possible involvement of AIF and Endo G. This unique pathway may address drug resistance issue which has become a hindrance for most conventional chemotherapeutics.

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0-47

Discovery of potent small molecule inhibitor for dengue through *in silico* and *in vitro* approaches

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Background

Dengue is a catastrophic arboviral disease, globally and locally. At present there are no effective antiviral drugs and only a limited approved vaccine available¹. The much-needed treatment for dengue has pushed studies to look into strategizing and developing methods to search for prospective inhibitor candidates. Due to the pivotal role of dengue NS2B/NS3 viral protease in the viral replication, the inhibition of this enzyme is considered to be the key strategy for the development of new dengue antiviral drugs². In this present study, we sought to identify and evaluate small molecules by adapting in silico and in vitro approaches in search of potent inhibitor against the dengue NS2B/NS3 protease.

Methodology

To that aim, total of 80 small molecules from the Riken NPDepo Authentic library were screened. Based on the highest percentage of inhibition exhibited, the hit compounds were chosen and molecular docking was performed to predict the binding model of the compounds to dengue NS2B/NS3 protease. Then, a chromogenic-based protease inhibition assay was carried out to evaluate whether the selected candidates inhibited dengue protease activity in vitro.

Results and Discussion

Based on *in silico* evaluation, computational simulation revealed the free energy of the compound I, II and III are comparable to that of quercetin, a known NS2B/NS3 protease inhibitor. Through the docking evaluation also showed that all three compounds bound to the DENV NS2B/NS3pro near the active site, hence confirming the possible interaction between the compounds and dengue NS2B/NS3pro. Interestingly, our results in the protease inhibition assay carried out indicated that compound I, II and III have the ability to inhibit dengue NS2B/NS3 proteolytic activity. **Conclusion**

The *in vitro* inhibition results of the three compounds that we obtained from this study, agrees well with the docking outcome, hence, portrays a potential inhibition activity against dengue virus.

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O-48

Systematic review on preclinical reports: Titania Nanotube Arrays technology for medical orthopaedic screw implant application
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Background

Titania Nanotube Arrays (TNA) technology is a promising surface technology for medical implants, especially for orthopedic applications. The long-term stability of a surface technology on a medical implant is frequently associated with osseointegration. This review provides an overview on TNA technology for medical orthopaedic screw implant applications from preclinical reported studies. **Methodology**

The review was conducted based on PRISMA-P protocol and predetermined keywords such as "orthopedics implants screw", "bone implant", "Orthopaedic Screw Implant" "TNA surface", "In vitro", "Ex vivo" "In vivo" "Titania Nanotube Array", "titanium dioxide nanotube arrays", "TiO2 nanotube arrays", "preclinical studies", "preclinical testing" and "osseointegration". Search engine database such as PubMed, Springer Link, Science Direct and Google Scholar were used. The pre-clinical studies published in English language within the past 5 years (2016 till 2021) were included. To broaden the search results, the terms were combined interchangeably by using the Boolena operators 'and' or 'or'. The titles and abstracts of each article were initially pre-screened. Studies involving those in predatory or blacklisted journals were excluded. To ensure transparency, replicability, and reanalysis feasibility, every step of this approach was me-

of all extracted research. Results and Discussion

The search has generally generated 778 articles. In further analysis, 30 articles were finalized according to inclusion and exclusion criteria. Data were further grouped into *in vitro, ex vivo, in vivo* studies. Most preclinical reports discussed the ideal TNA physiochemical properties that enhance cell proliferation, differentiation and surface properties that could contribute to long-term osseointegration activity.

ticulously recorded. To prevent double duplication, Endnote software version X7 was used as a reference manager to combine the results

Conclusion

The systematic reviews reported on TNA's ideal physiochemical properties are crucial for long-term bone-implant interaction. Knowledge from this study may contribute to the cutting-edge development of biomedical implant technology.

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0-49

Clinicopathological association of Chronic Rhinosinusitis with Nasal Polyp (CRSwNP) and periostin expression

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Background

In the recent years, periostin (POSTN), a gene encoding an extracellular matrix protein with similarity to fasciclin family has emerged as a potential biomarker for various types of cancers. Besides that, POSTN also plays a role in stimulating eosinophil migration and activation in chronic inflammation and immune pathways. Chronic rhinosinusitis (CRS) with nasal polyp (NP) (CRSwNP) is mediated by Th2-/eosinophilic inflammation type of immune response [1]. POSTN has been proposed to be found in NP tissues and contribute to formation of NP in CRS patients. In this study, we investigated the POSTN protein expression in NP tissue and then we determined its association with the clinicopathological features of CRSwNP patients.

Methodology

Tissue samples were collected from 24 CRSwNP patients and their clinicopathological features such as demographic data (age, gender, race and smoking history), clinical (history of bronchial asthma and/or atopy, SNOT-22 scores, Lund-Kennedy scores, Lund-Mackay scores), haematological (percentage of serum eosinophils and total IgE) and pathological features (tissue eosinophil count, degree of inflammation, mucosal ulceration, squamous metaplasia, and fibrosis) were evaluated. POSTN protein expression was assessed by immunohistochemical analysis. A single linear regression analysis was performed to find the association of POSTN protein expression with the clinicopathological features.

Results and Discussion

Expression of POSTN protein was detected in all 24 NP samples. The mean IHC score POSTN is 7.31. There was no significant association of POSTN protein expression with clinicopathological features (gender, smoking history, history of bronchial asthma

and/or atopy, SNOT-22 scores, Lund-Kennedy scores, Lund-Mackay scores, percentage of serum eosinophils, total IgE, tissue eosinophil count and other histopathological features (p value of 0.66, 0.11, 0.64, 0.57, 0.76, 0.17, 0.09, 0.21, 0.66, and 0.57 respectively) except for age (p=0.045). Our findings of POSTN expressed in all NP tissue staining the area of subepithelial tissue and infiltrating inflammatory cells was consistent with other study [1-2]. POSTN is produced in fibroblasts, endothelial and epithelial cells by interleukin (IL)-4 and IL-13 stimulation. Fibroblasts is found to be abundant in NP tissue. Tissue eosinophilia (eosinophil count >10 per high power field) is known to be a hallmark for CRSwNP and a predictor for NP recurrence. Ninomiya et al. 2018 found that protein expression associated with severity of CRSwNP but not in our study. Even though we excluded those patients taking steroids which can suppress the periostin production, we should have exclude those who was asthmatic and on any immunomodulator. Lebrikizumab was shown to be effective in controlling exacerbation of asthmatic patients with high POSTN level in their serum [1]. A larger sample size study at multi-centre setting would be more representative of CRSwNP population.

Conclusion

Although presence of POSTN is detected within all nasal polyp tissues, however, POSTN protein expression was not significantly associated with clinicopathological features of CRSwNP. Therefore, tissue POSTN appears to have no role in evaluation of patients with CRSwNP.

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O-50

Detection of herpes simplex virus-1 by direct immunofluorescence and viral isolation from cerebrospinal fluid

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Background

Herpes simplex virus (HSV) is the human herpesvirus that leads to herpes simplex encephalitis or meningoencephalitis and is frequently lethal if not treated properly. Here, we described a case of a 21-year-old man who presented with acute confusion and abnormal behaviour and was later diagnosed with HSV-1 meningoencephalitis based on immunofluorescence and viral isolation from cerebrospinal fluid.

Methodology

A lumbar puncture was performed immediately during admission to the ward. Cerebrospinal fluid was also sent for viral culture. The culture was inoculated into human cells in culture (HEp-2) cell monolayers and observed for cytopathogenic effect (CPE). Then the slide was prepared for direct immunofluorescence staining using fluorescein isothiocynate-conjugated HSV type 1 and HSV type 2 antisera. Positive findings would demonstrate cells with fluorescent staining, whereas negative specimens would demonstrate cells with a reddish-brown counterstain. Informed consent to publish had been obtained

Results and Discussion

Brain MRI was performed for further evaluation, which showed a focal area of gyral thickening at the left frontoparietal lobes with leptomeningeal enhancement at the left Sylvian fissure, suggestive of meningoencephalitis, and no hydrocephalus was noted. The CSF results revealed 0 polymorphs cells/mm³ and lymphocyte count, with 0 pus cells. CSF biochemistry showed glucose of 1.67 mg/dl and a very high total protein of 1596 mg/dl. The results of viral culture were obtained on admission day ten. After 10 days of culture with daily CPE observation, CPE evidence of HSV was detected. The prepared slide was observed under the ultraviolet microscope and revealed positive for HSV-1 and negative for HSV-2. HSV-1 has accounted for more than 90% of all herpes simplex encephalitis cases in adults and children. It spreads by oral contact and primarily results in cold sores, while HSV-2 is sexually transmitted and causes genital herpes.

Conclusion

It has been proven that immunofluorescence antigen detection is a quick, accurate, and sensitive method for distinguishing HSV-1 and HSV-2 antigen in the cerebrospinal fluid of those infected individuals.

0-51

Development of cisplatin-resistant urothelial cancer cells using pulse-shock treatment

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Background

Cisplatin has been used in chemotherapy of advanced urothelial carcinoma (UC) that is no longer amenable to local treatment such as surgery and radiotherapy. However, almost 30% of UC patients develop resistance to chemotherapy and recurrence within 2 years post-therapy. The understanding of the mechanism of cisplatin resistance in UC patients is scarce. In pulse-shock chemotherapy, cancer cells are subjected to chemodrug treatment in a short period of time.

Methodology

In this study, cisplatin-resistant UC model was developed and analysed for morphological and transcriptomic changes. Firstly, cisplatin IC50 dosage of a urothelial cancer cell line, 5637 cells was determined. Cells were then subjected to repeated pulse-shock treatment with the respective IC50 dosage and morphological changes will be recorded. At the end of the repeated treatment cycles, IC50 of the cisplatin-resistant cell was compared to its parental counterpart and transcriptomic changes in drug resistance genes were analysed.

Results and discussion

When pulse-shock treated for 2 hours, cisplatin IC50 dosage of 5637 cells was 11uM. After sixth treatment cycle, IC50 was significantly increased to 21uM, which was 2 times higher compared to parental cells. During the course of treatment, morphological changes vary from polyploid giant cells, MSC-like cell, parental cells and bizarre shaped cells. ABCA1, ABCG2, Bax and BIRC5 genes were found to be significantly upregulated in cisplatin-resistant 5637 compared to parental cells. Stress induction through pulse-shock cisplatin treatment promotes cancer plasticity through transformation into polyploid giant cells. These polyploid giant cells has been reported to drive cancer progression through activation of EMT, drug resistance, apoptotic inhibition and many more. Molecular mechanism and regulation of polyploid giant cells are not fully understood, hence, an exciting niche awaiting exploration. Conclusion

The model has a potential to be used for evaluation of existing and novel targets for recurrent therapeutic regimes, thus making UC management more efficient, convenience and increased patients' quality of life.

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0-56

Selection of ssDNA aptamers against Programmed Death-Ligand 1 (PD-L1)

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Background

Recently, serum PDL1 expression has been demonstrated to be a potential biomarker for cancer patient's treatment response and outcome or can even be used for early cancer detection. The most common biomarker detection approach utilizes antibody-based, but the use of antibody face numbers of drawback. Consequently, another potential alternative to antibody known as aptamers have become increasing popular. Aptamers exhibit characteristic that can overcome the drawbacks of antibody such as lower production cost, minimal batch-to-batch variation, possess better thermal stability and offers variety of chemical modification. Thus, this study aims to isolate ssDNA aptamers against recombinant human PD-L1 for future diagnostic purpose.

Methodology

In this study, we demonstrated the isolation of ssDNA aptamers against rhPD-L1 via agarose bead-based Systematic Evolution of Ligands by Exponential Enrichment (SELEX). The SELEX enrichment monitoring was determined by direct Enzyme-Linked Oligonucleotide Assay (ELONA). The enriched SELEX cycle was then cloned, followed by sanger sequencing to identify the sequence of potential aptamers. The sequences were analysed using MegaX software, online mFold software and online MEME (Multiple Em for Motif Elicitation).

Results and Discussion

Total of 9 SELEX cycle was performed and SELEX cycle 6 was selected as an enriched cycle. Pool of ssDNA from SELEX cycle six was cloned and a total of 36 positive colonies were sequenced. Phylogenetic analysis of the positive colonies yields 9 clusters distinct aptamers. Further analysis of Gibbs free energy, frequency of a sequence and significant motif of all the sequences revealed 3 potential aptamer sequence.

Conclusion

These 3 potential aptamer binding affinities should be characterized as it could be potential biorecognition element targeting the PD-L1 for the diagnostic application.

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P-1

Suppression of microRNA-9 promotes anti-tumour activity in KMS-28BM human multiple myeloma cells

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Background

Multiple myeloma (MM) is the second most common blood cancer characterised by clonal expansion of malignant plasma cells within the bone marrow [1]. Epigenetic aberrations such as microRNAs (miRNAs) dysregulation are of great significance in the development and malignancy of MM. MiRNAs are short, single stranded non-coding RNAs (~19-25 nucleotides) that play an important role in regulating post-transcriptional expression of target genes. Up-regulation of miR9 has been reported in various cancers including MM; however, the mechanisms underlying its aberrant expression and functional alterations in MM are still unclear [2].

Methodology

This study aims to investigate the functional role of miR-9 in MM cells *in vitro*. KMS-28BM MM cells were transfected with miRNA-9 inhibitors or miRNA inhibitors scrambled negative control. The expression levels of miR-9 vs control were determined by RT-qPCR. The effect of miRNA-9 suppression on cell proliferation was assessed by

MTS assay. Flow cytometry analysis using Annexin V-FITC/PI double staining was used to measure the proportion of apoptosis cells. **Results and Discussion**

The RT-qPCR results demonstrated that miR-9 inhibitors were successfully delivered and suppressed the miR-9 expression level in KMS-28BM compared to the control (*P*<0.05). Furthermore, MTS and flow cytometry analysis revealed that suppression of miR- 9 expression significantly decreased cell growth and increased the number of early apoptosis cells in KMS-28BM, respectively (*P*<0.05). Our previous microarray-based miRNA expression profiling results showed that miR-9 is upregulated in the majority of MM patients and cell lines, suggesting miR-9 plays a pivotal role, at least in part, in myelomagenesis. The present study findings further confirm that miR-9 functions as an oncogene in MM pathogenesis, and suppression or silencing of miR-9 inhibits cell

Conclusion

Suppression of miR-9 promotes anti-tumour activity by inhibiting proliferation and inducing apoptosis in MM cells indicates that miR-9 may be a potential therapeutic target in MM.

proliferation and induces apoptosis in KMS-28BM MM cells.

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P-2

Association of oxidative damage measured by 8-hydroxyguanosine formation with altered risks to hepatocellular carcinoma in Malaysian study population

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Background

Hepatocellular carcinoma (HCC) is the most devastating type of liver cancer. This disease is the third leading cause of cancer deaths with a 5-year survival rate of 7% [1]. Cancer is the pathogenesis product of DNA damage resulting from multiple factors which among others is oxidative stress. Oxidative stress can be detected by the DNA base damage, through the formation of 8-oxo-7, 8-dihydro-2'deoxyguanosine (8-oxodG) [2]. The 8-oxodG acts as oxidative stress indicator and has been essentially specified as a recognized initiator of the carcinogenic process and premutagenic injury in mammalian cells [3]. In this preliminary study, we investigated the possible association of oxidative DNA damage in a form of 8-oxodG in HCC patients in comparison with Malaysian healthy controls. We also analysed the effect of 8-oxodG in different races and gender in both groups.

Methodology

DNA of peripheral white blood cells was isolated from 91 HCC patients and 304 controls. The level of oxi dative DNA damage was determined by highly sensitive ELISA kit produced from Japan Institute for Control of Ageing (JalCA), Nikken Foods. Ltd. The determination of oxidative DNA level was digested by Ravanant (1998) and Evans (1999) methods with some modifications. The results were reported as 8-oxodG per 50 µL DNA sample in ng/ml.

Results and Discussion

Quantitative measurement of 8-oxodG was higher in HCC patients at mean value of 3.30 \pm 2.32 ng/ml. In controls, the average value is 1.57 \pm 1.92 ng/ml. There was a significant difference in the average value of 8oxodG level between the controls and HCC patients where p < 0.001. Comparison between gender showed that there was a significant difference observed in the level of 8-oxodG between male and female in controls (p < 0.05). The level of 8-oxodG was higher in male (1.736 \pm 2.033) ng/mL than in female controls (1.087 ± 1.433) ng/mL. From previous studies, it was reported that smoking, alcohol consumption and bad sleeping pattern were the sporadic factors for male tendency to get the oxidative damage [4]. Hepatocarcinogenesis has been reported to be associated with DNA damage through the process of oxidation. Oxidation in turn, leads to the huge amount of transversion from G

T in tumor genes such as p53 gene [5]. In this study, however, Malays and no-Malays showed no significant difference in their level of 8-oxodG indicating no disparity. Conclusion

HCC patients showed greater oxidative damage to DNA as compared to controls. This suggests that oxidative DNA damage may contribute to the pathogenesis of HCC. Since 8-oxodG was higher in males, it is indicating that the males are at a greater risk of developing HCC than the females.

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P-3

Mechanisms of optical light properties using optical spectroscopy for various honey detection

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Background

Honey is a natural sweetener made by honeybees from plant nectar that has been consumed by humans for thousands of years [1]. There are two kinds of honey which are monofloral honey and multifloral honey. Monofloral honey is made from a single plant nectar that has been transformed into honey. It includes Acacia, Sidr, and Kelulut honey. Meanwhile, multifloral honeys like Tualang, Manuka, and Kelulut honey are made from a blend of several different plant nectar types found in deep forests or mixed by the beekeeper before being converted into honey [2]. Honey consumption in Malaysia is increasing due to its various nutrient contents that are beneficial to one's health. Different types of honey contain varying amounts of sucrose, fructose, and glucose, which contribute to the sweetness of honey.

Methodology

In this study, pure Tualang, Kelulut, Acacia, Manuka, and Sidr honey samples were used. Optical spectroscopy in the ultraviolet-visible-near infra-red (UV-VIS-NIR) range was used to observe the optical properties of all of the honey, including absorbance and transmission. The wavelength range covered during the measurement was 350 nm to 1662 nm. The Ocean View software was used to capture the spectra of all honey. To improve the accuracy of the results, the spectra was collected 15 times for each honey. The average of the results were plotted, and boxplot analysis was performed for all spectra results.

Results and Discussion

The spectra results were taken based on the wavelength of the optical properties, which were absorbance and transmittance. Based on the observed results, the peak absorbance spectra for all honey in both UV-VIS-NIR were similar. Further analyses were done using boxplot analysis to differentiate the characteristics for all type of honey. Sidr honey has the highest mean absorbance value compared to other honeys. The current study found that all Malaysian raw honey met the international standard because the sum of their fructose and glucose contents were less than 60% [3]. But since Manuka and Sidr honey originated outside of Malaysia, it was assumed that both honeys do not adhere to the standard and thus consisted a higher sugar content compared to other honey. According to the Beer Lambert Law equation, increasing the sugar content will increase the number of molecules in the solution, which will increase the absorbance value. Based on previous research, an increasing pattern in absorbance value was highly noticeable as sugar concentration increased. Maximum transmission spectra for all honey in both ranges were close for both UV-VIS and VIS-NIR ranges. From the transmission boxplot analysis, Sidr honey recorded the lowest mean compared to other honeys. As a result, when the concentration of the molecules increased, the transmission value decreased, giving Sidr honey the lowest mean among the others.

Two optical properties, which are absorbance and transmission, were studied for all of the honey using optical spectroscopy in both the UV-VIS and VIS-NIR ranges. A boxplot analysis was performed on the spectra results to compare the mean value of the peak spectra. Sidr honey recorded the highest mean absorbance value compared to other honeys since it has the highest sugar content. According to Beer Lambert's law, as the concentrations of molecules increase, the absorbance value increases but the transmission value decreases. Therefore, Sidr honey has the lowest transmission mean value compared to other honeys.

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P-4

The effects of small molecules chaperone on cell viability in primary neonatal fibroblast cell line: Toxicity indicator for pharmacological chaperone therapy

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Background

Hunter syndrome (HS) is a lysosomal storage disorder (LSD) caused by a mutation in the gene *IDS*, which produces the defective enzyme iduronate-2-sulphatase (IDS). Due to the limitations of enzyme replacement therapy for the treatment of HS, the use of small molecule compounds as pharmacological chaperone therapy (PCT) has recently been widely investigated [1]. Usually, small molecules with higher binding affinity that can be used at low concentrations level will be chosen as pharmacological chaperone candidates [2]. As a result, *in vitro* assays such as methyl tetrazolium test (MTT) assay can be used to determine the cytotoxicity of small molecules before proceeding with *in vivo* studies. Therefore, the aim of this study is to investigate the cytotoxic effects of selected small molecules on the viability of primary neonatal fibroblast cell line.

Methodology

Selected small molecules of heparin tetrasaccharide, (H004), heparin octa-saccharide (H008), heparin decaoctasaccharide (H018), heparin disaccharide standard (HD001) and chondroitin dermatan trisulphate (CD007) were diluted in minimum essential medium (MEM) with several concentrations (0.16-500 µM). The BJ CRL-2522 fibroblast cell lines were seeded into 96-well cell culture plate at a density of 1x104 cells/well before the medium was removed from the cells after 24 hours. Subsequently, the cell lines were treated with the medium containing the respective small molecules for 72 hours (5% CO₂, 37oC). The cell viability was assessed by MTT assay, and the half maximal inhibitory concentration (IC₅₀) was calculated to determine the toxicity of small molecules.

Results and Discussion

The cell viability was decreased in a dose- and time-dependent manner when treated with the respective small molecules. HD001 has significantly exhibited concentration and time- dependent inhibitory effect to the cells with the highest IC $_{50}$ of 239 μ M. H018 showed IC $_{50}$ of 18.82 μ M making it the most toxic among all small molecules tested. The IC $_{50}$ of H004, H008 and CD007 were 83.52 μ M, 68.13 μ M and 72.03 μ M, respectively. Despite having non-toxic outcomes, HD001's chaperone action is inadequate to serve as a pharmacological chaperone because it binds to the IDS at a high concentration (inhibition constant, K; 431.4 μ M). [2]. On other hand, the high binding affinity of CD007 has a greater chaperone effect (K, 21 μ M) compared with HD001 with a compromised moderate toxicity level towards the cell lines.

Conclusion

This study concludes that CD007 could be considered as potential pharmacological chaperone for PCT in HS based on its safety profile.

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P-5

Decrease in mitochondrial dynamics in a patient with energy deficiency disorder

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Mitochondrial dynamics refers to the coordination of fission and fusion process in mitochondria for the maintenance of their distribution, shape and size. Mitochondrial dynamics is crucial in the regulation of cell cycle, immunity, apoptosis and mitochondrial quality control of various processes such as biogenesis and mitophagy. Defective mitochondrial dynamics due to mutations were reported to be linked with human diseases [1]. For example, patients with energy deficiency are suspected of having primary mitochondrial disorders that can be inherited through defects of either nuclear (autosomal-dominant, autosomal recessive, X-linked) or mitochondrial (mtDNA) genes [2]. In this study, we aim to characterize the mitochondrial dynamics of a patient with energy deficient disorder.

Methodology

Fibroblasts from a 2-month-old male patient with clinical symptoms of encephalopathy, seizure, cardiomyopathy, severe metabolic acidosis, pulmonary hypertension idiopathic and low level of lactate were cultured and harvested for western blot experiment. Fibroblast from American Type Culture Collection (BJ CRL-2522™) was used as normal control. Total protein was extracted, and the concentration was measured by Bradford assay. Next, samples were probed antibodies with GTPases of the dynamin superfamily. Antibodies used were Mitofusin 1 (Mfn1), Mfn2, Optic Atrophy 1 (Opa1) and Dynamin-related Protein 1 (Drp1). Anti-glyceraldehyde 3-phosphate dehydrogenase (GAPDH) monoclonal antibody was used as normalization control.

Results and Discussion

Mitochondrial fusion protein levels of Mfn1; Mfn2; and Opa1 were found to be significantly decreased in this patient's cells as compared to wild type (WT) (p<0.05). Meanwhile, there were no differences observed in mitochondrial fission protein level of Drp1 (p>0.05) as compared to WT cells. From these results, the reduced fusion proteins may contribute to energy deficiency in this patient. Previous studies suggested that mitochondrial fusion is known to allow functional complementation of mitochondrial DNA, protein and metabolites, whereas mitochondrial fission facilitates mitochondrial transport, mitophagy, and apoptosis [3].

Conclusion

Our study suggested a defective mitochondrial dynamic that appeared to affect the fusion process. Further studies such as immuno-fluorescent staining is highly recommended in order to gain complementary insights into mitochondrial dynamics as well as its involvement in many primary mitochondrial disorders.

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P-7

Selectivity and storage stability of the aptamer-modified multiwalled carbon nanotubes screen-printed carbon electrode for direct detection of 25-hydroxyvitamin D_2

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Background

Electrochemical biosensors are some of the promising platforms that are low cost, easy to use with the possibility of being integrated into portable devices for point-of-care use. Recently, aptamer-based electrochemical biosensors have received a significant attention because of their advantages such as good stability, high sensitivity and selectivity. The immobilization of the aptamer onto the working electrode is very important so that the target recognition by the aptamer can be transduced into a measurable electrical signal [1]. Here, we described the selectivity and storage stability of an aptamer-based electrochemical biosensors for direct detection of 25 (OH) vitamin D_2 on the aptamer-modified multi-walled carbon nanotubes screen-printed carbon electrode (MWCNTs-SPCE).

Methodology

Aptamer specific to 25 (OH) vitamin D_2 was immobilized onto the working electrode surface of the MWCNTs-SPCE. The performance of the modified electrodes was observed to determine their selectivity towards 25 (OH) vitamin D_2 . The long-term storage stability of the aptamer-modified MWCNTs-SPCE was investigated by storing the fabricated devices for a period of up to 30 days at 4°C. Electrochemical measurements were carried out using electrochemical impedance spectroscopy (EIS) in a redox probe (0.1 M KCl containing 5.0 mM Fe (CN) $_6^{3-/4-}$).

Results and Discussion

The immobilization of the aptamer was determined through the differences of resistance to charge transfer (Rct) values obtained. The Rct values increased when aptamer was immobilized onto the surface of MWCNTs-SPCE compared to the bare MWCNTs-SPCE. The specific reaction between aptamer and 25 (OH) vitamin D_2 causing the Rct value to be decreased. However, there were no changes observed when the assay carried out in the absence of the aptamer. This demonstrates that the signals were due to the specific interaction and not to non-specific adsorptions [2,3]. A good linear response was observed for different 25 (OH) vitamin D_2 concentrations in the range of 0.1 - 30 ng/ml with a detection limit of 0.72 ng/ml. For the long-term storage stability, the aptamer-modified MWCNTs-SPCE showed a consistent response for the first 14 days, then experiences a drop afterwards. Therefore, the aptamer-modified MWCNTs-SPCE was determined to be stable for 14 days at 4°C .

The aptamer-modified MWCNTs-SPCE was shown to be selective towards 25 (OH) vitamin D_2 and had a reasonable shelf-life of around 2 weeks which can be further improved for a better long-term storage.

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P-

Spectrum of F9 mutations in Malaysian Haemophilia B patients

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Haemophilia B (HB) is a rare X-linked bleeding disorder characterized by deficiency in coagulation factor IX (FIX) due to mutation within Factor 9 (F9) gene [1]. F9 gene is located at Xq27.1-q27.2, consists of 8 exons and 7 introns, dispersed across 34 kb length [2]. Severity of HB patients can be classified into mild, moderate and severe based on the coagulation FIX level. Patients with FIX level <1% are classified as severe whereas patients with FIX level 1%-5% and >5% to <40% are classified as moderate and mild, respectively [3]. In Malaysia, there are limited studies on F9 gene mutations. Thus, this study aimed to identify the mutations within a representative cohort of HB patients in Malaysian population.

Methodology

A total of 41 non-familial related HB patients were studied. The patients' blood samples and clinical details which contained information on factor level, disease severity and inhibitor status were obtained from National Blood Centre and hospitals all over Malaysia. Genomic DNA was extracted from the blood samples using QlAamp® DNA Blood Midi Kit (Qiagen) and subjected to polymerase chain reaction (PCR). Eight different primer sets were used to amplify exon/intron borders and all eight exons of F9 gene. PCR products were then sequenced using ABI 3730XL DNA analyzer. Sequencing data of patients were aligned against reference sequences, NG_007994 and NM_000133.4 using CLC Main Workbench to identify the mutation and deduce amino acid changes.

Results and Discussion

F9 gene mutations were successfully detected in all patients, which comprise 32 point mutations (26 missense, 5 nonsense and 1 silent), 6 deletions, 1 insertion, 1 insertion-deletion (indel) and 1 splice site mutation. These mutations were distributed throughout all exons, except exon 3 with most of the mutations located in exon 8 (~42%). Five of the discovered mutations in Malaysian HB patients, c.874C>T, c.523_539delinsGAA, c.318_319insGGC, c.230T>G and C.40delC were found to be novel, whereas the remaining variants have been reported by various countries in several online Haemophilia B databases. In our study, two patients developed inhibitor against FIX and the mutations detected in these patients are nonsense mutation (c.1150C>T) and deletion (c.40delC).

Conclusion

Our study identified heterogenous mutation profile in HB patients in Malaysian population. These findings are important for genetic confirmation of HB patients, genetic counselling, prediction of inhibitor development and also for carrier screening among family members.

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P-9

Small RNA signatures from isoniazid-resistant *Mycobacterium* tuberculosis

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Background

Tuberculosis (TB) caused by *Mycobacterium tuberculosis* (MTB), remains a significant public health concern worldwide. The emergence of multidrug-resistant TB (MDR-TB) (resistance to at least isoniazid or rifampicin) has significantly jeopardized the treatment and control of TB. Isoniazid-resistant TB (Hr-TB) is much more common than rifampicin-resistant TB (RR-TB) [1]. The regulatory mechanism underlying the emergence of Hr-TB remain to be fully elucidated. Small non-coding RNAs (sRNAs) regulate bacteria essential functions such as survivability and metabolism, among others. Also, the sRNA has been implicated as one of the caused for development of drug resistance [2]. The present study investigates the expression profile of sRNAs in drug-susceptible TB (DS-TB) and Hr-TB strains using next generation sequencing (NGS) with Illumina HiSeq2500 Deep Sequencing Technology for better understanding of the mechanism of resistance to isoniazid in MTB.

Methodology

Three DS-TB (SBH49, SBH149, and SBH372) and three Hr-TB strains (SBH365, SBH438, and SBH509) isolated from pulmonary TB patients were cultured in BD BACTECTM MGITTM (Becton Dickinson, USA) and their drug susceptibility to first-line antibiotics, i.e., streptomycin, isoniazid, rifampin, and ethambutol were tested with BACTECTM MGITTM 960 SIRE kit (Becton Dickinson, USA). Resazurin microtiter assay (REMA) was performed to determine the minimum inhibition concentration (MIC) for isoniazid. Next, RNA from both susceptible and resistant strains were extracted using MasterpureTM Complete DNA and RNA purification kit (Lucigen, USA). The integrity of RNA was determined by Agilent RNA 6000 Nano kit (Agilent Technologies, USA). Small RNA libraries preparation was carried out using NEBNext® Small RNA Library Preparation kit (NEB, UK). Samples were sequenced by single-end reads of 50 base pairs generated through Illumina HiSeq 2500TM in Rapid Run mode. RNA sequences were trimmed for adapter sequence and low-quality reads were filtered out. The sequences were annotated to reference genome (MTB H37Rv) and differential analysis were carried out with CLC Bio Genomic Workbench (version 8). sRNAs target prediction was performed by using TargetRNA2.

Results and Discussion

DS-TB strains were susceptible to all first-line antibiotics, while Hr-TB strains only resistance to isoniazid with MIC of 0.125µg/mL for SBH365, and 2mg/mL for SBH 438 and SBH509. A total of 63,252,209, 63,636,812, and 61,148,224 qualified Illumina reads were obtained from DS-TB strains and 75,296,115, 58,965,389, and 67,935,496 reads from Hr-TB strains. The overall *de novo* assembly of sRNA sequence data generated 255, 255, and 255 for all DS-TB strains and 310, 359, and 403 for Hr-TB strains with sRNA length of 18-30bp. Comparative analysis revealed that 728 sRNAs were differentially expressed in the Hr-TB compared with the DS-TB, of which 422 were upregulated and 306 were downregulated. sRNA target prediction using computational method reveals these sRNAs target wide range of mRNAs which involve in toxin production, survivability, and others [3].

Conclusion

sRNAs play a major role in the regulation of gene expression and mediates the cellular processes in bacteria. This data demonstrates that sRNA may serve as an invaluable resource for revealing the molecular basis of the regulation of expression associated with the mechanism of isoniazid resistance in MTB.

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P-11

Determining an optimal DNA isolation method for rodent fecal samples by 16S rDNA bacterial diversity identification

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Background

The choice of an optimum DNA isolation method from fecal samples is a constant challenge as ineffective lyses of diverse bacterial cells may lead to bias detection in the representation of a bacterial community in a sample. In addition, degraded DNA and presence of PCR inhibitors such as humic acid and polysaccharides carried over from fecal samples have been known to further reduce PCR efficiency. The purpose of this study is to determine an optimal DNA extraction method for rodent samples by assessing the general bacterial diversity based on PCR products amplified from the partial 16S rDNA gene. The PCR-DGGE technique separates the identical size of PCR amplicons by their differential mobility on gel based on sequence variations.

Methodology

In this study, five (5) fecal samples from wild rodents were collected and added into the Zymo Research DNA/RNA Shield reagent as preservative. Six DNA extraction methods i.e. QIAamp PowerFecal Pro DNA Kit (QIAGEN), the QIAamp AllPrep PowerViral DNA/RNA Kit (QIAGEN), the QIAamp UCP Pathogen Kit (QIAGEN), the ZymoBIOMICS DNA Miniprep Kit (ZYMO Research), and two conventional methods using different component in lysis buffer which were guanidium thiocyanate and CTAB were tested using between 10 to 15 mg of feces. The samples were extracted according to manufacturer's protocol and previous article [1] with some modifications, respectively. The extracted DNA were then subjected to amplification of V2 to V3 region of the bacterial 16S rDNA gene and DGGE separation of the amplicons. The DGGE banding pattern was analysed using the GelCompar II software (Applied Maths, Belgium) to compare the microbial diversity by using the Shannon-Weaver index [2].

Results and Discussion

The results showed that DNA yield varied with the extraction method; where the conventional method using quanidium thiocyanate showed a higher yield (average 324.22 ng/ul) than the other methods (average 45.33 ng/ul for the QIAamp PowerFecal Pro DNA Kit, 51.92 ng/ul for the QIAamp AllPrep PowerViral DNA/RNA Kit, 220.74 ng/ul for the QIAamp UCP Pathogen Kit, 42.4 ng/ul for Zymo-BIOMICS DNA Miniprep Kit, and 165.74 ng/ul for the conventional method using CTAB). Majority of DNA extracted showed degradation when checked by gel electrophoresis. The QIAamp PowerFecal Pro DNA Kit was observed to show intact DNA. The DGGE profiles showed that the QIAamp PowerFecal Pro DNA Kit and QIAamp All-Prep PowerViral DNA/RNA Kit extracted the highest number of DNA bands (total of 168 and 167), while QIAamp UCP Pathogen Kit, Zymo-BIOMICS DNA Miniprep Kit, and both conventional methods using guanidium thiocyanate and CTAB recorded 161, 160, 147, and 152 total of bands, respectively.

Conclusion

As a conclusion, QIAamp PowerFecal Pro DNA kit is the optimum DNA isolation method for rodent fecal samples as it provides better quality of DNA and microbial diversity in their DGGE profiles. Furthermore, this method is much time efficient QIAamp PowerFecal Pro DNA Kit and QIAamp AllPrep PowerViral DNA/RNA Kit produced more bands on their DGGE profiles than the other methos due to their use of bead-containing lysing matrix and vigorous homogenization.

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P-12

The toxic effects of p-Cresyl Sulfate on bone metabolism

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Background

p-Cresyl Sulfate (pCS) is a uremic toxin that has been implicated in kidney disease, cardiovascular risks, endothelial dysfunction and neuropathies. In chronic kidney disease (CKD), pCS progressively accumulates in the body as the dysfunctional kidneys have a reduced ability to excrete toxins normally. pCS accumulated as a consequence of CKD cannot be removed from th body through dialysis, hence leading to further accumulation. This ultimately leads to bone loss which correlates with the worsening of CKD. As such, pCS could play a part in the development of bone loss with CKD. The objective of this review is to further understand the comprehensive effects pCS has on the bone.

Methodology

An extensive literature search was conducted in PubMed using the following keywords, 'p-Cresyl Sulfate' OR 'p-Cresol Sulfate' OR 'p-Cresyl Sulphate', 'uremic toxin' OR 'uraemic toxin' AND 'bone' OR 'osteoblast' OR 'osteoclast' OR 'osteocyte'. From 2013 to 2022, 54 papers were found that contained the following keywords. Out of the 54 papers, 27 papers with significance were selected. The inclusion criteria for the study are in vivo and in vitro studies that examined the effects of pCS on bone. The exclusion criteria for the study is as follows: review article. After reviewing each article, 5 papers were selected for this review.

Results and Discussion

pCS is a prototype protein-bound uremic toxin associated with a multitude of toxic biological and biochemical effects. pCS is a substrate for human organic anion transporters (hOAT) 1 and 3 which is expressed in osteoblasts and other tissues. hOAT1 and hOAT3 may have a physiological role as large-capacity pCS transporters which could be a factor in the accumulation of pCS to toxic levels in osteoblasts [1]. Accumulation of pCS affects sclerostin production. As an inhibitor of the Wnt-signalling pathway, sclerostin prevents osteoblast formation and osteoprotegerin (OPG) production, inhibits osteoblast-mediated bone formation, and stimulates bone resorption by stimulating RANKL expression in osteocytes [2]. pCS also increase apoptosis and reduce osteoblast proliferation and viability with or without increased oxidative stress. This manifests by downregulation of the parathyroid hormone receptor (PTHR) on osteoblasts and a decrease in PTH-stimulated cAMP production through activation of c-Jun N-kinase (JNK) and p38 mitogen-activated protein kinase (MAPK) [3]. and triggers JNK phosphorylation by attenuating PTH responsiveness in the bones under CKD conditions.

Conclusion

This review evaluates the toxic effects pCS induces on bone. However, further studies must be conducted to understand the mechanism of action and full effects pCS has on bone metabolism. In the future, pCS could potentially become a new therapeutic target for the management of bone disorders.

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P-13

Characterization of ssDNA aptamers against ACE2 protein as therapeutic targets for COVID-19 infection

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Background

Coronavirus disease 2019 (COVID-19), a highly contagious and rapidly spreading disease with significant fatality in the elderly population has swept across the world for the past three years since 2019. The enormity scale of this pandemic has resulted in the emergence of several SARS-CoV-2 variants with Omicron, the current main circulating variant of concern [1]. Nonetheless, all the variants identified to date share the same cell entry mechanism where the process is initiated when the spike protein of the viruses attach to the angiotensinconverting enzyme 2 (ACE2) receptors on their host cells [2]. Therefore, blocking the spike protein-ACE2 interaction using a biological binder targeting the ACE2 is a viable strategy for COVID-19 treatment. Aptamer is a short length of nucleic acid [RNA or singlestranded DNA (ssDNA)] which is selected through an in vitro selection called systematic evolution of ligands by exponential enrichment (SELEX). The selected aptamers have a 3D conformation that can specifically bind to their target with high affinity and pose many superior properties such as being smaller in size, thermally stable, and nearly non-immunogenic.

Methodology

Single-stranded DNA library of 76-bp oligonucleotides containing a randomized core region of 40 nucleotides, flanked by primer binding regions of 18 nucleotides on each side (5'-ATACCAGCTTATTCAATT-N40-AGATAGTAAGTGCAATCT-3') was synthesized. Recombinant Histagged ACE2 protein (BBI Life Sciences, China) was immobilized to Ni-NTA Magnetic Beads (Gold Biotechnology, USA). Aptamers which showed high affinity towards the ACE2 were then selected from the initial library using SELEX. The isolated aptamers were cloned using the PCR Cloning Kit (NEB, USA) and the plasmids were extracted using DNA-spin Plasmid DNA Extraction Kit (iNtRON Biotechnology, Korea). The plasmids were sequenced and the resulting aptamer sequences were subjected to the UNAFOLD web server (http://www.unafold.org/) for ssDNA secondary structure prediction, followed by

RNA tertiary structure prediction using RNAComposer (https://rnacomposer.cs.put.poznan.pl/), and finally, converted to equivalent ssDNA tertiary structure using Discovery Studio Visualizer v3.5. The aptamer-ACE2 interaction was predicted using the HDOCK docking program (http://hdock.phys.hust.edu.cn/).

Results and Discussion

Aptamers specifically bound to ACE2 were isolated after 13 rounds of magnetic bead-based SELEX. Here, we reported one of the isolated aptamers, Apt15, which potentially binds to the ACE2. The UNAFOLD revealed that this aptamer has a hairpin loop structure and single-stranded region with a Gibbs free energy value of -0.10. The low free energy value indicates that the ssDNA structure is thermodynamically stable [3]. *In silico* 3D molecular docking demonstrated that the single-stranded region of Apt15 binds to the ACE2 at the site recognized by the SARS-CoV-2 spike protein with a confidence score of 0.9894, thus suggesting the potential application of this aptamer as a therapeutic target for COVID-19 infection.

Conclusion

In conclusion, one ssDNA aptamer targeting ACE2 with therapeutic potential against COVID-19 was successfully identified in this study. Further investigations are necessary to determine its binding affinity to ACE2 and its ability to block the virus spike protein-ACE2 interaction. In future, this aptamer could serve as a broad-spectrum inhibitor against any existing or future emerging viruses that also use ACE2 for cell entry.

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P-14

Detection of Campylobacter jejuni and Campylobacter coli from retails broiler chicken by duplex PCR

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Background

Campylobacter is one of the leading causes of foodborne diarrhoea illness in the developed countries and cause a significant public health concern worldwide. This study was designed to determine the prevalence of Campylobacter coli and C.jejuni contamination in poultry retails meat in Kota Bharu by duplex PCR from direct samples.

Methodology

A total of 50 fresh and chilled poultry chicken meat were purchased from 13 retail markets in. The samples were put into polyethylene bag and wash with Phosphate-buffered solution, the bacterial lysate was prepared directly from chicken wash, enriched in CCDA broth for 48 hours and the presence of *C.coli* and *C.jejuni* were detected by duplex PCR.

Result and discussion

Overall results revealed 22 % contamination of $\it Campylobacter$ occur in poultry retail meat in Kota Bharu, Kelantan. Out of the fifty

samples of fresh (n= 16) and frozen/chilled (n=34) chicken thigh, 10/ 50 (20%) was positive for C.coli and/or C.jejuni. Majority (9/10, 90%) of contamination were C.coli whereas, the remaining (1 sample) was C. jejuni. Frozen/chilled samples have lower frequency of Campylobacter contamination,11.8% (4/34) as compared to fresh samples, 37.5% (6/16). No C.jejuni contamination was observed in fresh samples, and only 2.9% (1/34) in frozen/chilled samples. While C.coli detected at rate 6/16(37.5%) in fresh chicken samples and 4/34(11.7%) in frozen/chilled chicken samples. The prevalence of Campylobacter spp in retail meat Kota Bharu was 22% (11/50) of fresh and frozen chicken meat is relatively low as compared to other result reported previously in Japan and Egypt who obtained retail level rate varying from 64-68%. Meanwhile, the reported prevalence of Campylobacter spp in broiler chicken in Malaysia even greater up to 97.1%, other studies done in East-cost Malaysia shows 20% of contamination from total of 120 cloaca swabs taken whereas the later obtain 45%-70%. The contamination attributed by the cross-contamination of poultry carcasses during defeathering, evisceration and carcass chillers, which reflected by the contamination rate of cases and meat. The result showed higher prevalence of contamination with C.coli and C.jejuni in fresh chicken samples compared to frozen/chilled. Previous studies showed the influence of chilled meat and inclusion of skin as determining factors in the prevalence of Campylobacter spp. Out of 34 chilled chicken samples, 11.8 % (4/34) were positive with Campylobacter spp, this 3-fold higher in fresh chicken samples, 37.5% (6/ 16). This in consistent with study done in central part of Malaysia, stated that apart from chilled temperature, packing in polyvinylidene film-overwrapped storiform trays may indirectly creating microaerophilic conditions which facilitate the survivability of Campylobacter.

The prevalence of *C.coli* and *C.jejuni* among poultry retails chicken meat in Kota Bharu, Kelantan is low as compared to limited data in other part of Malaysia. However, more studies with a larger sample sizes and involvement of retail market are required.

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P-1

The efficacy and safety of PD-1/PD-L1 immunotherapy in endometrial cancer: a meta-analysis

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Background

The programmed cell death protein 1 (PD-1)/Programmed cell death ligand 1 (PD-L1) pathway has a crucial role in the immune escape mechanism and growth of cancer cells in various cancer types, including endometrial cancer [1]. In exploring better treatment options, immune checkpoint inhibitor studies of PD-1/PD-L1 in endometrial cancer are actively conducted in clinical trials with promising findings in other cancers. However, the efficacy of those immunotherapy remains inconclusive, and guidelines are inconsistent [2]. Therefore, this meta-analysis aims to provide an updated and robust analysis of the effectiveness and safety of PD-1/PDL1 immunotherapy in endometrial cancer on the objective response rate (ORR), disease control rate (DCR), and adverse events.

Methodology

A literature search was conducted using the database of PubMed, Scopus, and Web of Science databases by using the keywords of "endometrial cancer" OR "PD-1 inhibitor" OR "PD-L1 inhibitor" OR "immunotherapy" OR "immune checkpoint inhibitor" OR "atezolizumab" OR "avelumab" OR

"dostarlimab" OR "durvalumab" OR "nivolumab" OR "pembrolizumab". Meta-analysis of proportion and association was pooled using STATA version17 and RevMan version 5.4 software, respectively. The effect of immunotherapy on outcome parameters was estimated by effect size (ES) or odds ratio (OR) with 95% confidence intervals (Cls) for each study.

Results and Discussion

Five studies between 2017 and 2022 met the inclusion criteria, with 480 endometrial cancer cases undergoing PD-1/PD-L1 immunotherapy clinical trials. The pooled proportion of ORR undergo PD-1/PDL-1 inhibitor treatment was 26.47% [95% CI = 16.68-37.52]. Subgroup meta-analysis showed the pooled ORR of the proficient mismatch repair (pMMR) group was 8.80% [95% CI = 0.20-24.39], and which was 37.03% [95% CI = 23.22-51.95] of the deficient mismatch repair (dMMR) group. Metaanalysis proportion of DCR undergo PD-1/PD-L1 inhibitor treatment was 41.44% [95% CI = 36.38-46.60]. Subgroup meta-analysis showed the pooled proportion DCR of the pMMR group was 24.97% [95% CI = 14.87-36.49] and dMMR group was 45.02% [95% CI = 39.26-50.85]. The efficacy of the PD-1/PD-L1 inhibitor showed a significantly higher ORR in dMMR than pMMR [OR = 6.30; 95% CI = 3.60-11.03]. Results were the same for DCR, a significantly higher in dMMR than pMMR [OR = 2.57; 95% CI = 1.66–3.99]. The safety pooled proportion of patients with at least one adverse event was 69.19% [95% CI = 62.56-75.46], and grade three or higher facing adverse events was 15.38% [95% CI = 12.01-19.04]. Based on the MSI status subgroup analysis of ORR and DCR, the efficacy of PD-1/PD-L1 immunotherapy in endometrial cancer was significantly better in dMMR patients.

Conclusion

PD-1/PD-L1 immunotherapy in endometrial cancer shows a promising outcome in early clinical trials, found to have better efficacy in the dMMR population. Thus, patients with dMMR are more suitable for this treatment. Further studies on targeted immunotherapy approaches need to be fully explored before it can be applied for a better treatment outcome.

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P-17

Assessment of the NGS data using FastQC for quality control in the study of antiproliferative effects of dichloromethane *Clinacanthus nutans* fractions extracts on human breast cancer cell lines for biomarker discovery

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Background

In Malaysia, Clinacanthus nutans (CN) leaves are traditionally used as a treatment for breast cancer. However, scientific approval of the mode of action of the CN extract on MCF-7 cell lines at the molecular level is still lacking. High throughput Next-Generation Sequencing technology (NGS) to obtain large sequence numbers that benefit genetic medical research was used in this experiment [1, 2]. Bioinformatics tools such as FastQC and Picard were used to evaluate sequence data for quality control as an essential method

during the analytical pipeline [3] that is important to identify challenges of systematic bias challenge in this study. The objective of this study was to evaluate the quality of NGS data sequence using FastQC in the early part of transcriptomic analysis pipelines of antiproliferative effects of dichloromethane CN fractions extracts on human breast cancer cell lines for biomarker discovery.

Methodology

Dichloromethane CN leaves extract with 50% inhibitory concentration (IC₅₀) value of 108µg/mL was exposed to the human breast cancer cells, MCF-7 for 72hrs chosen from a previous study. RNA was extracted from the treated and untreated cells for transcriptomic sequence, NGS technology using the Illumina HiSeq4000 platform. The sequencing output was the raw data that must be removed from the adapter sequences by the short read trimmed using FastQC (Babraham Bioinformatics, UK). This analysis uses the Linux operating system, Ubuntu 18.0v and the adapter was removed by a barcode tool, FlexbarFlexbarth its command line and script. All results were presented in multiQC which aggregated the results and generated a single HTML report with plots to visualize and compare various metrics between the samples involved.

Results and Discussion

The output of multiQC was filed from FastQC. In the summary report of FastQC status, important attention was made based on sequence quality and sequence length distribution. After RNA sequencing using the paired ends modules samples generated GC overall percentage of 54% and a length of 150bp showing no difference in GC composition and no biased library complexity, differences in amplification, or library specific causes. The sequence quality resulted in a pass or good result in the green area and more than 30 phred scores for all samples. As a result, the total overrepresented sequences found in each library make up more than 0.1% of the total that was passed and good quality which does not affect the biological consideration. Overall, the quality of RNA-seq data was high and the sequence quality was also good.

Conclusion

Multi QC report using FastQC is an advantageous tool because it is relatively quick to generate and provides a clear method for comparing samples to determine consistency and identify problematic samples. From this study, the multiQC report contains high quality clean data that was downstream in all future transcriptomic analyses for biomarker discovery in breast cancer research.

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P-18

IGF-1 and IGFBP-1 expressions as the potential prognostic biomarkers in women with Endometrioid Endometrial Cancer (EEC)

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Background

Insulin-like growth factor-1 (IGF-1) and the IGF binding protein 1 (IGFBP-1) expressions have been shown to play a vital role in cancer biology, including endometrioid endometrial cancer (EEC). We examined the prognostic value of these locally expressed biomarkers in endometrial biopsies and correlated them with clinicopathological data of EEC.

Methodology

mRNA expression of *IGF-1* and *IGFBP-1* in the endometrial biopsies were analysed in patients with EEC (n=25) and control (n=25) cases using quantitative polymerase chain reaction (qPCR) method. These data were then validated using immunohistochemistry (IHC) analysis and combined with EEC cases form a separate cohort (n=71) comprised of consecutive patients who underwent hysterectomy at UKMMC, between the year of 2014 to 2019. Overall survival was evaluated using the Kaplan-Meier method, with differences compared using the log- rank test. Independent relationships between these biomarkers and clinicopathological data were assessed using multivariate logistic regression models.

Results and Discussion

The IGF-1 and IGFBP-1 mRNA expressions were not significantly different between both groups, EEC vs. control. However, IGF-1 expression in IHC analysis was observed to be highly expressed in the EEC compared to the control group, while IGFBP-1 had a low expression in the EEC cases (P<0.05). IGF-1 was significantly associated with prognostic features of EEC (P<0.05), while no associations were found in the IGFBP-1 expression. In our sub-analysis, high IGF-1 and negative IGFBP-1 expression were significantly correlated with poor progression-free survival (PFS) in advanced stage of EEC (P<0.05). Univariate and multivariate analyses showed that IGF-1 served as a predictive biomarker in EEC survival. Therefore, we postulate the continuous high expression of local IGF-1 protein in EEC cells could lead to poor outcomes. Our findings support that the circulating estrogen and IGF-1 were independently associated with a higher risk of recurrence in patients with stage III and IV of EC (Merritt et al. 2021). We also propose that a shift in the local expression of IGFBP-1 and IGF-1 may serve as a biomarker for the prognosis of EEC development.

Conclusion

Local expression of IGF-1 and IGFBP-1 may serve as prognostic biomarkers for EEC.

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P-19

Pharmacological consequence of inhibiting MAPK p38 using Ralimetinib dimesylate on lipopolysaccharide-induced E-selectin and VCAM-1 expression in HUVEC

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Background

Endothelial dysfunction plays a prominent role in the pathogenesis of sepsis and is associated with life-threatening organ dysfunction. Lipopolysaccharide (LPS), a Gram-negative bacterial component, is an important sepsis-associated mediator that induces the expression of adhesion molecules such as E-selectin and VCAM-1 upon its binding to dedicated pattern recognition receptors on endothelial cells (EC) [1]. Endothelial E-selectin and VCAM-1 expression promotes leukocyte adhesion, and high leukocyte infiltration in various organs is associated with a poor prognosis in sepsis patients. As the expression of E-selectin and VCAM-1 is partly driven by the

activation of the MAPK p38 signaling pathway [2], it is unknown whether Ralimetinib dimesylate (RM), a selective p38 MAPK inhibitor, can be used as a treatment to reduce E-selectin and VCAM-1 expression once LPS-driven activation of EC has started. RM treatment was previously shown to reduce TNF- α production in LPS-induced macrophages *in vitro* [3]. In this study, I investigated the pharmacological effect of RM treatment on E-selectin and VCAM-1 expression in LPS-stimulated HUVEC was investigated.

Methodology Ten μ M of RM was added into the HUVEC medium at 0.5, 1, 1.5, and 2 hours after HUVEC was exposed to 1 μ g/ml of LPS. After 2, 4, 5, and 6 hours of LPS exposure, the cells were trypsinized and subjected to flow cytometry analyses. The Mean Fluorescence Intensity (MFI) of E-selectin and VCAM-1- conjugated fluorochromes were determined in the treatment groups and statistically compared to LPS-stimulated HUVEC controls using one-way ANOVA and Bonferroni post-hoc test. Each group was represented by three biological replicates. The results were reported as mean + SD. Viability of HUVEC was monitored microscopically.

Results and Discussion

LPS induced the expression of E-selectin and VCAM-1 in HUVEC in a time-dependent manner. E-selectin and VCAM-1 were maximally expressed on HUVEC at 4 and 6 hours after LPS exposure, respectively. E-selectin expression was attenuated throughout the 6 hours' duration of LPS exposure upon post-LPS treatment with RM at 0.5, 1, 1.5, and 2 hours after LPS exposure. VCAM-1 expression was not affected upon post-LPS treatments with RM. These findings suggest that MAPK p38 to be an important signaling cascade mediating LPS-induced E-selectin expression and can be pharmacologically targeted to attenuate E-selectin, but not VCAM-1 expression.

Conclusion

RM can be used to pharmacologically target the p38 MAPK signaling pathway to attenuate the expression of E-selectin, but not VCAM-1, in LPS-activated EC. Follow-up *in vivo* study should be done to investigate the effect of RM on the expression of E-selectin and VCAM-1 in animal model of experimental sepsis.

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P-20

AST, ALT, Bilirubin and AST/ALT Ratio Role; COVID-19 Patients

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Background

Impaired liver function upon admission has been linked to the severity of COVID-19 infection, yet the data is debated [1]. Therefore, this retrospective study aimed to evaluate the liver function among COVID-19 patients during hospitalization and its association with the disease severity.

Methodology

Patients aged 18 to 80 years with positive COVID-19 at Hospital Raja Perempuan Zainab II (HRPZ II), Kota Bharu, Kelantan, with available AST, ALT, Bilirubin, and AST/ALT ratio data on admission, were retrospectively evaluated from March 2021 until March 2022. Disease severity was categorized based on the Annex 2e guidelines by Ministry of Health Malaysia, which further classified them into mild to moderate disease (Stage 1-3) and severe to critical illness (Stage 4-5). The AST, ALT, Bilirubin, and AST/ALT ratio levels on Day 1 admission were archived from the electronic medical record system and compared between the two groups. Statistical analysis was performed using SPSS version 27. This study was approved by (JEPeM-USM) with protocol code USM/JEPeM/21100691 and the Ministry of Health Malaysia NMRR-21-762-58458 (IIR).

Results and Discussion

The study involved a total of 168 COVID-19 patients with a mean (SD) age of 46.67(16.10) for mild to moderate and 56.66(12.41) for severe to critical. There was a significant age group for both groups (pvalue=0.002). During hospitalization, 16(14.41%) patients progressed to death from severe to critically ill patients. Upon admission, the median (IQR) of AST and ALT were significantly higher in the severe to critical group compared to the mild to moderate group, [AST; 39.0(49.0) and 24.0(14.0), ALT 38.0(43.0) and 21.0(18.0)], p<0.05. However, there were no significant differences between both groups for bilirubin level and AST/ALT ratio. Non-survivors had higher AST and ALT levels compared to survivors, with median (IQR) of [AST 98.0(88.0) and 32.0 (26.0), ALT of 67.5(90.0) and 28.0(31.0), (p<0.05). Similarly, there were no significant differences between nonsurvivors and survivors for bilirubin and AST/ALT ratio. Our study supports the idea that abnormal liver function at admission has been shown to be associated with the disease severity and mortality of COVID-19 infection. Therefore, there is a need to observe hepatobiliary sequelae in COVID-19 survivors as there are dynamic changes in liver function following hospital discharge.

Conclusion

Abnormal AST and ALT level at admission has been shown to be associated with the disease severity and mortality of COVID-19 infection. Further study needed to evaluate liver damage in COVID-19 post-discharge.

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P-21

MALDI-TOF mass spectrometry-based strategy in discovery reproducible N-glycans in human fibroblast

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Mass spectrometry (MS) has evolved over the years into a powerful tool that facilitates the identification of glycoproteins and *N*-glycans. *N*-glycans are essential for controlling metabolic pathways and cellular communication between cells, tissues, and organs. Because of this, there is a lot of interest in the study of glycosylation and *N*-glycans as biomarkers for several diseases [1]. Thus, the reproducible release and identification of *N*-glycans from tissue are important criteria for biomarker discovery. The aim of this study is to explore a general out-look on *N*-glycans reproducibility in fibroblast using matrix assisted laser desorption ionization time-of-flight (MALDI/TOF) mass spectrometry.

Methodology

Two different batches (batch 1 and 2) of the human fibroblast cell lines were grown to confluence subjected to FANGS protocol [2] and permethylated before mass spectrometry analysis. Sample from these two batches (Batch 1: A1, B1 and C1; Batch 2: A2, B2 and C2) were processed on different days and the released permethylated glycans were analyzed on different days using MALDI/TOF. Triplicate spots from each sample were spotted on a MALDI plate and analyzed three times. The resulting [M+Na] ⁺ glycan intensities from the spectra corresponding to the same sample were averaged and used to prepare bar charts.

Results and Discussion

Results showed that there were no significant differences between different spots of the same sample in all samples (p>0.05). This finding is similar to the recently published paper [3], in which the plasma *N*-glycan analysis of control samples was used to evaluate the reproducibility of sample processing. Reproducibility of sample processing on different days showed a few sialylated glycans were missing in A2 and B2 compared to A1 and B1 samples. This is possibly due to the low levels of the collected sialylated glycans thatmay be lost during sample processing or possibly due to biological variation during cell growth. There were differences in the signal intensities of the Hex5HexNAc2 and NeuAcFucHex5HexNAc4 species between the two batches. Most of the signal intensities are similar in the two batches prepared from the C cells.

Conclusion

In conclusion, this technique of *N*-glycan identification from fibroblast using mass spectrometry can be applied in investigations using human fibroblasts. Further features must be improved such as to reduce the variability of *N*-glycans obtained during sample processing or cell growth.

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P-22

Spray-dried andrographolide nanoparticles for the treatment of Alzheimer's disease: Preparation and cytotoxicity evaluation

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Background

Andrographolide is a major bioactive constituent found in *Andrographis paniculate* (Hempedu bumi in Malay). It has been reported to possess a promising neuroprotective effect that has the potential for the treatment of Alzheimer's disease [1]. However, the penetration of andrographolide into the central nervous system via the blood-brain barrier is one of the main obstacles for the treatment of Alzheimer's

disease. Oral administration of *A. paniculata* extract has been reported to limit the reach of andrographolide to the brain owing to high plasma protein binding (55%) [2-3]. Andrographolide also readily degraded in acidic and alkaline environments of the gastrointestinal tract. Hence, it is hypothesised that if andrographolide in the nanoparticle form is delivered to the brain, the poor bioavailability limitations could be overcome. The current study aimed to produce spray-dried andrographolide nanoparticles and investigate their cytotoxicity in vitro using a human neuroblastoma cell line.

Methodology

Andrographolide nanoparticles were produced using a nano-spray dryer (Buchi Nano Spray Dryer B-90 HP). The particle size was measured using laser diffraction technology and electron microscopy was used to observe the morphology of the particles. The cytotoxicity of the nanoparticles was tested on SH-SY5Y neuroblastoma cells. The nanoparticle suspension at various concentrations was added into the cells in two-fold dilutions and incubated for 48 hrs. 0.05% resazurin was then added and further incubated for 4 hrs. Fluorescence readings were taken at a wavelength of 590nm following excitation at 544nm using a microplate reader. The half-maximal inhibitory concentration (IC₅₀) was determined by comparing the inhibition to the untreated cells. In addition, lactate dehydrogenase (LDH) assay, nitrite oxide (NO) level and reactive oxygen species (ROS) level were determined using the supernatant of the cells following the assay kit guidelines. An andrographolide solution was included as a control. Three independent experiments were performed in triplicate.

Results and Discussion

The spray-dried andrographolide nanoparticles were spherical with a rough surface with a hydrodynamic diameter of 724 \pm 39 nm with a polydispersity index of 0.3. The spray-dried nanoparticles showed lower toxicity (IC50 of 15.22 \pm 0.61 $\mu g/ml$) as compared to the solubilised andrographolide control (IC50 of 10.82 \pm 0.61 $\mu g/ml$). As anticipated, cells treated with solubilised andrographolide showed a higher amount of LDH release compared to the formulated nanoparticles, indicating that the formulation slightly reduced the toxicity of the andrographolide. The nitrate level in the supernatant did not increase dramatically compared to the untreated control, indicating that the cellular death did not involve the release of NO level in the cells. Similarly, the cells did not show any differences in ROS between the treated and untreated cells, suggesting that cellular death might not be induced through the ROS pathway.

Conclusion

These findings warrant further in vivo studies to establish the efficacy of the spray-dried andrographolide nanoparticles for the treatment of Alzheimer disease.

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P-23

Cross-reactivity between Leptospira Serovars in microscopic agglutination test for Leptospirosis

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Leptospirosis is a zoonotic disease resulting in high morbidity and mortality in Malaysia and other endemic countries. Microscopic agglutination test (MAT) is the gold standard method in diagnostic leptospira armamentarium. MAT is a laborious serological test performed by observing agglutination reaction of patient sera when incubated against a panel of live Leptospira serovars. This study aimed to analyze the degree of cross-reactivity of sera for MAT conducted in Institute for Medical Research (IMR).

Methodology

The study included all sera for MAT in 2019. MAT was performed using an extended WHO panel comprising 20 serovars and 4 strains of *Leptospira species*. A titer of more than or equal to 1:400 is considered seropositive. Data analysis for serological cross reactivity was performed using Microsoft Excel for descriptive statistics.

Results and Discussion

96 sera (2.4%) out of 3965 sera were tested positive for MAT. Single seropositivity was observed in 40.6% of the sera and the remaining demonstrated cross-reactivity ranging from 2 to 9 serovars (average of 3 serovars). The high degree of cross reactivity observed may be a unique interplay between acute febrile illness (AFI) sera reacted to an extended MAT panel. Single serovar seropositivity involving serovar Fugis, Whitcombi and Lai Langkawi, *L. meyeri, L. borgpetersenii* serovar Batavae, may reflect a convalescent serum reacting to specific infecting serovar. Our finding is in parallel with other studies that support the optimization of extended MAT panels to suit locally dynamic circulating serovar. The limited number of paired sera in this study hinders the optimal determination of important infecting serovars.

Conclusion

An extended MAT panel tailored for local endemic serovar displays a high degree of cross-reactivity in AFI sera that serves as a more sensitive tool for the diagnosis of leptospirosis.

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P-24

Susceptibility of human lung-derived cell lines MRC-5 and A549 to Malaysian SARS-CoV-2 isolate

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Background

Severe Acute Respiratory Disease 2 (SARS-CoV-2) has been identified as the causal factor to the recent COVID-19 pandemic [1]. One of the capacities required to study this virus is to isolate and propagate them according to the requirement of the studies. To date, the majority of studies utilized monkey-derived Vero

cells to rapidly propagate the virus. However, due to its highly susceptibility to SARS-CoV-2 infection and limited capacity to metabolize drugs, this cell line is not suitable for pathophysiology and anti-viral research. Vero cells are not preferred for investigation of pathological mechanism of host cell's response to virus infection because they are not derived from human lung tissue. Furthermore, they are difficult to be used in the study to assess cytopathic effects as they do not express type 1 interferon genes [2]. In this study, we adapted the SARS-CoV-2 isolates into selected human lung-derived cell lines, i.e., MRC-5 and A549, to investigate the capacity of these cell lines as a platform for the characterization of SARS-CoV-2 isolated in Malaysia. MRC-5 has already been identified to be highly susceptible to the infection of various human coronaviruses, including HCoV-OC43, HCoV-229E and Middle East respiratory syndrome coronavirus (MERS-CoV) [3]. Meanwhile, A549 was selected due to its origin of lung derived.

Methodology

To determine the growth profile of SARS-CoV-2 in human-derived cell lines, a local SARS-CoV-2 clinical isolate obtained from the IMR archive was first adapted into Vero E6, thus resulting in the generation of passage 1 (P1). The P1 isolate was used to study the growth profile of the virus in MRC-5 and A549 cell lines. P1 isolate was subjected to Whole Genome Sequencing (WGS) to identify the genomic sequence of the isolate. The replication kinetics of these isolates in MRC-5 and A459 were evaluated based on the on the formation of cytopathic effect (CPE), Cq value, plaque forming unit (pfu) as well as observation by electron microscope (EM). Whole Genome Sequencing (WGS) was repeated on the propagated SARS-CoV-2 passages to examine the genomic stability.

Results and Discussion

There was no formation of CPE observed after inoculation of SARS-CoV-2 into MRC-5 and A549 cells after 7 days of culture. In parallel, the qRT-PCR results suggested that the virus did not multiply well in these cells. Plaque assay and EM images further supported these findings. In terms of genome stability, WGS data revealed several genetic polymorphisms in the genome of the virus adapted in MRC-5 and A549 cells.

Conclusion

MRC-5 and A549 are not susceptible to SARS-CoV-2 infection.

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P-27

Effects of kaempferol on cell morphology, migration and microtubule functions in colorectal cancer cell HT-29

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Kaempferol, a naturally-occurring flavonoid, exhibits similar structure to estrogen and exerts estrogen-like activity. Extensive studies have been done to investigate its anticancer properties against various cancers [1,2]. Here we report effects of kaempferol on cellular activities that are closely regulated by microtubules. We also reveal our finding on expression of main microtubules components following kaempferol treatment in colorectal cancer cell line HT-29.

Methodology

We carried out cell viability assay to examine whether kaempferol inhibits HT-29 cell growth by estimating viable cell numbers. Next, we performed cell spreading and scratch wound assay to investigate the effects of kaempferol on HT-29 cell morphology and migration activity. To obtain a clearer understanding of how kaempferol causes alterations in microtubule network and cell motility, we performed whole cell microtubule analysis by flow cytometry.

Results and Discussion

Kaempferol decreased viable HT-29 cell numbers in a dosedependent manner with almost 66% decrease in the viable cell number within 24 hours after the addition of 240 µM kaempferol. During cell spreading, HT-29 cells treated with IC_{50} of kaempferol (143 μM) were found to be significantly reduced in cell area and increased circularity. This suggests kaempferol inhibits cell spreading and promotes a more circular-shaped cells. Further investigation on microtubule organization revealed that kaempferol suppressed normal microtubule spreading in HT-29 cells, consistent with reduction in the cell area. Kaempferol-treated cells were having intense antitubulin staining around the cell edge almost identical to the effects seen in taxol-treated HT-29 cells. Taxol treatment resulted in a more intense anti-tubulin staining, in agreement with its well-known role in promoting microtubule stabilization. More detailed microtubule analysis revealed that expression of α -tubulin and β -tubulin in kaempferol-treated cells was significantly lesser than untreated cells. Given the important roles of microtubule cytoskeleton in controlling cellular shape and cell motility, altered cell morphology and inhibition of cell migration found in kaempferol-treated HT-29 cells could be attributed to perturbation in microtubule functions.

Conclusion

Collectively, our findings show that kaempferol leads to alteration in cell spreading and hinders cell migration of cancer cells. These changes could be associated with decrease expression of α -tubulin and β -tubulin. Further work will be required to elucidate underlying mechanisms of kaempferol effects on microtubule dynamics.

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P-28

Genomic characterization of the first case of extensively drugresistant, travel-related *Salmonella enterica* Serovar Typhi in Malaysia

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Background

Typhoid fever, caused by *Salmonella enterica* serovar Typhi (S. Typhi) remains a serious global health concern. The effectiveness of antimicrobial therapy has been threatened by the emergence of Extensively Drug-resistant (XDR) S. Typhi exhibiting resistance to the first line and second-line antibiotic options. The aim of this study was to characterize an XDR S. Typhi isolate from a foreign worker with a history of recent travelling to Pakistan.

Methodology

Whole-genome sequencing (WGS) was performed on the isolate with Illumina MiSeq and data was analysed using Resfinder tool to detect the presence of genes associated with antimicrobial resistant profile. Pathogenwatch was used to characterize the genomic clonality in relation to the XDR outbreak of *S.* Typhi.

Results and Discussion

WGS analysis detected 10 antimicrobial resistance genes including aminoglycoside resistant gene (aph(3")-lb, aph(6)-ld, aac(6')-laa), trimethoprim resistant gene (dfrA7), sulphonamide resistant gene (sul1, sul2), beta-lactam resistant gene (blaCTX-M-15, blaTEM-1B), phenicol resistant gene (catA1) and quinolone resistant gene (qnrS). Similar gene profiles were seen for the XDR 5. Typhi from Pakistan. A core genome phylogeny tree constructed using selected genomes from Pathogenwatch collection clustered Malaysia XDR 5. Typhi isolate with XDR isolates from Pakistan, United Kingdom and Italy.

Conclusion

This first case of typhoid fever due to XDR *S*. Typhi detected in Malaysia highlights the need to be vigilant for future cases. Genomic characterization of *S*. Typhi using WGS as a tool in surveillance program is important to manage outbreak investigation by studying the pathogen population structure, identification and tracking of potential source both local as well as global.

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P-29

Suggestive SNPs of dengue pathogenesis towards severity in single-centre case-control study

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Background

Dengue has persistently served as a major public health concern especially in the tropical and sub-tropical areas. In Malaysia itself, dengue is a hyperendemic disease and the incidence rate continued to increase exponentially since the past ten years [1]. Manifestations of the dengue symptoms can range from asymptomatic, mild or severe. Dengue fever that progresses to severe dengue, either dengue haemorrhagic fever (DHF) or dengue shock syndrome (DSS), are usually life-threatening. The mechanisms of how this disease progresses to severity is incompletely understood. Early identification of cases

that are likely to progress to worse health conditions remains challenging. Multiple studies have highlighted the role of host genetics as a crucial factor towards severity. Thus, this study aimed to identify single nucleotide polymorphism(s) that predisposes dengue patients to severe disease outcomes among the Malaysian dengue cohort.

Methodology

This is a case-control study that includes a total of 188 dengue patients; 86 with dengue fever and 102 with severe dengue. Patients were recruited from Hospital Kuala Lumpur from year 2018 till 2020. Classification into dengue fever (DF) or severe dengue (SD) were made based on the WHO 2009 classification. DNA extracted from whole blood samples, were genotyped using Infinium™ Asian Screening Array. Single variant association analysis was performed on 617,245 single nucleotide polymorphisms (SNPs). Both sample- and SNP-levels data were subjected to stringent quality control checking for high quality genotype data.

Results and Discussion

We found genetic variations that are suggestively associated with dengue severity, namely rs9872672 and rs148681490. These SNPs reached the p-values of ~10-6, with their odds ratio of 4.574 (rs9872672) and 0.092 (rs148681490) were mapped to chromosome 3p14.2 and 11p11.2, respectively. Our findings at position 3p14.2 indicate a change of allele from C to T that occurred at a frequency of 0.143 globally [2]. Suggestive allele at this position identified an intronic region of the synaptoporin (SYNPR) gene that is predicted to be an integral component of synaptic vesicle membrane, whose probable function is linked with vesicular channel protein. Likewise, suggestive SNP found at chromosome 11 showed a change from A to G. The alleles at 11p11.2 encode for tetratricopeptide repeat domain 17 (TTC17) protein that are involved in the actin filament polymerization and cilium organization.

Conclusion

In conclusion, the findings from this study support the notion that host genetic factor contributes to the disease pathogenicity of dengue severity. Although the potential roles of the suggestive SNPs are still uncertain, further validation is highly warranted.

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P-30

Probiotics Alter Biofilm Formation of Periodontal Pathogen, Porphyromonas qinqivalis virulence-associated genes

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Background

Probiotics, a group of beneficial live microorganisms are suggested as an adjunct therapy to control the colonization of periodontal pathogens. Periodontal pathogens, especially *Porphyromonas gingivalis*, can form biofilm with other pathogens in the oral cavity and exhibit virulent factors that are detrimental to human gum health [1]. This study was carried out to assess the anti-biofilm activity of probiotics isolated from local fermented food and to investigate the underlying molecular activity behind the anti-biofilm activity.

Methodology

Cell-free supernatant (CFS) was prepared by centrifugation and filter sterilisation of the overnight culture of *Lactobacillus plantarum* FT 5. The probiotics culture was isolated from local fermented food and is kept as a private collection [2]. Anti-biofilm assay was performed against *P. gingivalis* where the optical density of *P. gingivalis* biofilm stained with crystal violet solution in the treated and untreated groups was measured. Sterile MRS and 0.2% chlorhexidine acted as negative and positive control respectively. Then, RT-qPCR procedure involving *fimA*, *mfa1*, *kgp*, and *rgp* genes was carried out on the treated and untreated *P. gingivalis*. Additionally, the 16S rRNA gene was employed as the normalization gene.

Results and Discussion

The anti-biofilm assay showed a significant reduction of *P. gingivalis* biofilm with a 90.5% (p-value < 0.05) reduction at the highest concentration of *L. plantarum* FT 5 CFS. The RT-qPCR revealed that biofilm reduction might be related to the downregulation of two biofilm-related genes, *fimA* and *mfa1*. Interestingly, *P. gingivalis* treated with *L. plantarum* FT 5 CFS showed upregulation of *kgp* and *rgp* genes. The results implicate that the inhibition of *P. gingivalis* biofilm formation by *L. plantarum* FT 5 CFS are due to the downregulation of fimbriae-related genes, *fimA* and *mfa1* genes. Aside from being vital virulent genes, *kgp* and *rgp* also contribute to the maturation of fimbrial protein [3]. Thus, the upregulation of *kgp* and *rgp* in *P. gingivalis* is probably necessary to overcome the downregulation of fimbriae-related genes.

Conclusion

In conclusion, *L. plantarum* FT 5 isolated from fermented foods can reduce the biofilm formation by *P. gingivalis* and the underlying mechanism might be related to biofilm and virulent-related genes regulation.

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P-31

Survival analysis of TNBC patients: A retrospective study from Hospital Universiti Sains Malaysia (HUSM)

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Background

Triple-negative breast cancer (TNBC), characterised by the absence of ER/PR expression with lack of HER2 overexpression, has drawn more attention both clinically and experimentally due to its aggressive biological characteristics and lack of effective treatment methods. There

are very limited population-based breast cancer survival and prognostic factors studies for Malaysian population, particularly that focus on TNBC patients. Hence, the purpose of this study was to determine the survival of TNBC patients at HUSM and to identify potential prognostic factors for their overall survival (OS).

Methodology

This retrospective study was conducted using primary data of 50 women diagnosed with malignant breast cancer at HUSM between January 2017 and December 2019 (USM/JEPeM/18120775). The patients were divided into two groups: TNBC and non-TNBC. Overall survival curves were estimated using the Kaplan-Meier method and a log-rank test was used to compare the survival functions among the two groups. Simple Cox's proportional hazard regression model was employed to identify the important prognostic factors of death.

Results and Discussion

The OS for TNBC was significantly lower than that for non-TNBC, at 45.0% vs 83.3% (p = 0.003). The mean survival time for TNBC was 38.7 months compared to 73.4 months for non-TNBC. Molecular subtype was a significant prognostic factor for malignant breast cancer patients whereby diagnosis with non- TNBC reduces the death hazard by a factor of 0.23 or 77.0% (95% CI 0.079–0.664; p = 0.007). Only staging status was statistically significant for OS of TNBC patients. The OS for early-stage TNBC was significantly higher than that for advanced-stage TNBC, at 66.7% vs 27.3% (p = 0.021). The median survival time was 57.0 months (95% CI 50.5-63.5 months) for earlystage TNBC vs 24.0 months (95% CI 19.7-28.3 months) for advanced stage TNBC patients. Advanced stage TNBC patients had 5.3 times increased hazard to death compared to early-stage TNBC patients (95% CI 1.096–25.647; p = 0.038). Since no other variables showed statistical significance on OS of TNBC or non- TNBC patients, multivariate analysis was not conducted. The OS and important prognostic factors reported in this study are in accordance with other studies [1,2] that have recorded worse OS for TNBC patients diagnosed at advanced stage. Nonetheless, other factors including patient's age, surgical treatment methods and family history of malignancy have been shown to be important prognostic factors for breast cancer [1,2], which was not the case for this study.

Conclusion

At HUSM, TNBC patients, particularly those diagnosed at advanced stage, have worse prognosis compared to non-TNBC patients. Future studies with larger sample size and comparative studies with other hospitals in Malaysia are necessary to enable the identification of more potential prognostic factors for TNBC and non-TNBC patients, with the aim to improve breast cancer management in this region.

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P-32

The shark VNAR single-domain antibody as potential binder for SARS-CoV-2

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Background

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) as an emergent zoonotic virus since December 2019, has caused coronavirus disease 2019 (COVID-19) as the first coronavirus pandemic in history [1]. Neutralizing antibodies against SARS-CoV-2 is useful, however with the large size (~150 kilodaltons (kDa)), complex hetero-tetrameric structure and susceptibility to extreme ambient temperature for the conventional antibody may limit their performance in therapeutic and diagnostic applications [2]. Antibody surface display technology allows the exploration of antibody fragments from other organisms, including VNAR (variable domain of immunoglobulin new antigen receptor (IgNAR)) from shark with better thermostability [3]. The advantages of shark VNAR single-domain antibody are due to possessing smaller size (~12 kDa) and simpler structure, with its antigen-binding capability comparable to conventional antibody [2, 3]. The aim of this study is to evaluate the potential of shark VNAR as binders towards SARS-CoV-2 as target antigen.

Methodology

A proprietary phage-displayed semi-synthetic shark VNAR library was subjected to 3 rounds of biopanning. The commercial wild-type SARS-CoV-2 receptor binding domain (RBD) diluted in various concentration was used as target antigen in this study. The biopanning steps per round involved immobilizing antigen on immunotube, blocking immunotube with skimmed milk, incubating phages from antibody library in immunotube, washing off unbound phages while eluting bound phages, followed by the re-propagation of specific phage clones in *Escherichia coli* TG1. The phage particles harvested from culture supernatant were applied in the next round of biopanning. Polyclonal phage pool and monoclonal phage clones from each biopanning round were analysed by agarose gel electrophoresis, to detect the presence of VNAR insert.

Results and Discussion

Throughout rounds of biopanning, the amplification of phage clones was observed in Round 3, with 11-fold enrichment as compared to Round 1. Based on agarose gel electrophoresis result, the presence of VNAR insert increased from 60% (Round 1) to 100% (Round 3), indicated that SARS-CoV-2 RBD-targeted phage clones have been enriched throughout subsequent rounds of biopanning. The identification and functional characterization of potential clones are in process.

Conclusion

Shark VNAR has been proven as potential binders targeting towards SARS-CoV-2. Thus, further biological characterization is required to determine the specificity and sensitivity of the potential binders.

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P-3

Selecting antibacterial aptamers against an essential outer membrane protein in *Pseudomonas aeruginosa* via aptamer repurposing approach

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Antimicrobial resistance is one of the major threats to global health, resulting in an increasing number of people suffering from severe illness or dying due to infections that were once easily curable with antibiotics. Pseudomonas aeruginosa represents one of the most concerning pathogens involved in antibiotic resistance where the World Health Organisation has classified this gram-negative bacterium as an ESKAPE organism, and it is also listed in the Priority 1: Critical list. Hence, in this project, we opted for a novel intervention by using aptamers to inhibit the growth of Pseudomonas aeruginosa, which can be cost-effective and less laborious. Besides, antimicrobial agents targeting the outer membrane protein (OMP) in Gram-negative bacteria can also be effective at killing or inhibiting bacterial growth as these proteins play an important role in bacteria survival as well as being exposed to the surface, which facilitates direct binding. Aptamer repurposing also reduces the cost and length of new drug development.

Methodology

Unmodified functional DNA aptamers from the Aptagen database were docked with the essential OMP using various docking tools such as HADDOCK 2.4 and HDOCK web server to determine the binding site and binding score of the various aptamers. Aptamers that bind to the active site with a good binding score were synthesised and folded into their 3D structures. *P. aeruginosa* cells were incubated with the aptamers overnight and the inhibitory effects on the bacterial growth curve were investigated.

Results and Discussion

Apt31 had the best HADDOCK score, and was found to bind near the active site of the OMP. Interestingly, this aptamer also exhibited significant antibacterial activity from the early stationary growth phase onwards. The addition of the aptamer after the late log phase also exhibited a similar effect. Besides, the antibacterial activity of apt31 was also dose-dependent. Hence, we can deduce that apt31 binds to the active site region and blocks the OMP activity. The major role of this OMP is to fold and insert β -barrel proteins into the outer membrane layer. As apt31 hinders the OMP activity, we postulate that the bacteria cells eventually die due to a lack of OMPs, which are essential for survival and virulence.

Conclusion

We show that the apt31, an aptamer that binds to an antitumor, is capable of binding to OMP in *P.aeruginosa*. The aptamer-OMP complex exhibits a significant antibacterial effect in a dose-dependent manner. Future experiments will include the expression of the recombinant OMP and determination of the dissociation constant (K_d) of the aptamer-OMP complex.

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P-35

Production of recombinant shark V_{NAR} single domain antibody specific against DENV Type 2 NS1

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Background

Dengue virus (DENV)) infection is one of the main global health issues leading to high morbidity and death rate. The accuracy of early diagnosis is the key for effective treatment. However, early diagnosis for DENV infection is a great challenge due to the similar onset symptoms of Dengue with other flaviviruses, concurrent infection

and cross-reactivity among dengue serotypes and other flaviviruses [1]. Besides, the limitations of conventional monoclonal antibodies (mAbs) also affect the efficiency of diagnostic kits [2]. Alternatively, the variable new antigen receptor (V_{NAR}) found in shark, has recently been identified as a promising diagnostic tool due to its excellent thermostability and capability to target on cryptic epitopes [3]. In this study, a potential clone that recognizing DENV Type 2 NS1 has been isolated and expressed. This recombinant protein was determined to be soluble and potentially developed as a novel reagent for DENV immunodiagnostic platform.

Methodology

A semi-synthetic library of shark V_{NAR} was used for antibody selection. Total of 3 rounds of biopanning was undertaken in respect to isolate potential binders which can specifically recognize DENV Type 2 NS1. The eluted phage from each round of biopanning were subject to polyclonal and monoclonal phage ELISA. After verified through DNA sequencing, the selected clone was expressed using bacteria expression system, followed by immobilized metal affinity chromatography (IMAC) purification process. The biological functions of recombinant V_{NAR} will then be characterized, including thermostability, specificity and sensitivity towards DENV Type 2 NS1.

Results and Discussion

For monoclonal phage ELISA, G3, Z3 and Z8 clones were identified to possess good affinity towards NS1 and lower cross reaction towards BSA. Thus, Z8 clone was selected to express as a 12 kDa protein with some optimization during protein expression. To verify target protein, the recombinant antibody reacted with anti-His6 antibody was performed using Western Blot. The functional assays of recombinant protein Z8 are currently underway.

Conclusion

Z8 clone isolated from biopanning has been identified as a potent binder. The recombinant anti-NS1 V_{NAR} Z8 antibody was successfully produced in a bacteria expression system. This new binder can be developed as new reagent for DENV immunodiagnostic platform.

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P-37

A fatal case of cerebral malaria complicated with *Gemella bergeri* bacteremia: Role of *Plasmodium* mitochondrial *cox3* gene and MALDI-TOF Mass Spectrometry for species identification

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Here, we described a fatal case of a 37-year-old returning traveller from Burkina Faso, West Africa, who presented with an acute fitting episode later diagnosed as cerebral malaria induced by *Plasmodium falciparum* based on microscopic examination and *Plasmodium* mitochondrial cytochrome c oxidase III (cox3) gene PCR target. Unfortunately, the patient passed away due to severe malaria with multiorgan failure complicated with *Gemella bergeri* bacteremia. *G. bergeri* was identified using MALDI-TOF mass spectrometry.

Methodology

Preliminary evaluation of blood film malaria parasite showed coinfection of *P. falciparum* and *Plasmodium malariae*. Further PCR-based detection analysis using the *Plasmodium* mitochondrial *cox3* gene was used to identify the malaria species. The patient's blood culture revealed gram-positive cocci in clusters, and further identification using MALDI-TOF Mass Spectrometry was done. Informed consent to publish had been obtained.

Results and Discussion

The patient blood sample was positive for *P. falciparum* based on analysis using a PCR assay targeting the *cox3* gene. Further, analysis done at Makmal Kesihatan Awam Kebangsaan confirmed *P. falciparum* infection. His blood culture revealed gram-shown in our case, cerebral malaria induced by *P. falciparum* causes coma, long-term positive cocci in clusters, which was later identified as *G. bergeri* using MALDI-TOF mass spectrometry. Despite the initiation of intravenous artesunate and subsequent parasite count responding and showing parasite clearance, the complication is detrimental. As neurological consequences and death. The patient also had concomitant fatal *G. bergeri* bacteremia. Initial examination revealed that the patient had poor oral hygiene and a missing tooth which could be the source of *Gemella* infection, which is common in the oral cavity and a causative organism of septic shock.

Conclusion

This report is the first fatal case of cerebral malaria caused by *P. fal-ciparum* coinfection with *G. bergeri* bacteremia. Therefore, it is critical to determine the exact aetiology for appropriate medical management. *Plasmodium* mitochondrial *cox3* gene and MALDI-TOF Mass Spectrometry are useful for species identification, as illustrated in this case.

P-38

Association of MTHFR polymorphism in stroke risk and rehabilitation outcomes

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Background

There are 12.2 million new stroke cases annually and about 101 million people worldwide are living with stroke aftermath [1]. In Malaysia, stroke is the third leading cause of death. The rs1801133 single nucleotide polymorphism (SNP) in the methylenetetrahydrofolate reductase (MTHFR) gene has been linked to stroke pathogenesis in a recent meta-analysis [2]. To date, there is a lack of study addressing this SNP towards stroke risk and rehabilitation outcomes in the Malaysian population. Hence, this study aims to investigate the

association of MTHFR rs1801133 SNP with the risk and rehabilitation outcomes in Malaysian stroke patients.

Methodology

The peripheral blood sample was collected from 251 age-matched individuals (113 stroke patients and 138 healthy controls without a stroke history) with written consent. The genomic DNA was extracted from these blood samples and the rs1801133 SNP was genotyped using a TaqMan® assay approach. The odds ratio and 95% confidence interval were calculated for risk association analysis. The stroke patients were subjected to a special-designed rehabilitation exercise and their pre- and post-exercise assessments including the Barthel index, Fugl-Meyer assessment-upper extremity, and Fugl-Meyer assessment-lower extremity were compared based on their genotypes.

Results and Discussion

This study showed that the presence of a T allele of the rs1801133 SNP had at least a 0.4-fold reduced risk of stroke. This suggests that the T allele of the rs1801133 SNP is protective against a stroke in the Malaysian population. Surprisingly, the finding of this study is in contrast with those reported in the meta-analyses [2,3]. Out of the 31 stroke patients who had completed their special-designed rehabilitation exercises, patients with a homozygous (C/C) genotype showed significant improvement in the post-rehabilitation assessments including the Barthel index, Fugl-Meyer assessment-upper extremity, and Fugl-Meyer assessment-lower extremity.

Conclusion

In conclusion, this study suggests that the MTHFR rs1801133 SNP is a potential biomarker for stroke risk and rehabilitation outcome predictions in Malaysian stroke patients. Data of this study could be useful for stroke management in the country.

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P-39

RNU6B and miR-16 as stable normalisation control for relative RT-PCR of Urinary microRNA in patients with colorectal polyps

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Background

Relative real-time polymerase chain reaction (RT-PCR) is an experimental technique widely used in quantifying the expression of a particular gene of interest. However, the reliability of gene expression data obtained through this technique is highly dependent on the stability of the housekeeping genes being used. Housekeeping genes act as a normaliser or endogenous reference in the relative

quantification of a target gene and its expression may vary between different types and sources of samples. Studies on urinary microRNA as a potential disease biomarker are expanding, and the suitable housekeeping genes are yet to be confirmed. In this study, we assessed the stability of three commonly used references for microRNA i.e., hsa-miR-26b, hsa-miR-16, and RNU6B.

Methodology

This study was registered (NMRR-19-1188-47227), approved by the Medical Review and Ethics Committee (MREC) of the Ministry of Health Malaysia, and the Human Research Ethics Committee of Universiti Sains Malaysia (JEPeM, USM), and conducted following the Good Clinical Practice Guideline. The urine samples were collected from 24 patients (12 with colonic polyps and 12 without colonic polyps) scheduled for colonoscopy at Hospital Sultanah Bahiyah, Kedah, Malaysia. Colonic polyps' diagnosis was confirmed by colonoscopy and histopathological report. MicroRNA was extracted from the urine supernatant based on the acid guanidinium thiocyanatephenol-chloroform-potassium acetate-lithium chloride method described by Zununi Vahed et al. (2016) with a slight modification. Reverse transcription and primer design for RT-PCR were performed following protocol by Balcells et al. (2011) and Busk (2014), respectively. We performed RT-PCR with cycling conditions of initial denaturation (95°C, 5 min) followed by 45 cycles of denaturation (95°C, 15 sec) and annealing (60°C, 30 sec). The analysis was extended with melting curve analysis (60°C to 95°C) to assess RT-PCR efficiency. The stability of Ct values was evaluated using three algorithms i.e., Best-Keeper, DeltaCq, and NormFinder. T-test was used to analyse whether a significant difference in mean Ct values of hsa-miR-26b, hsa-miR-16, and RNU6B existed between those two groups of patients.

Results and Discussion

There is no significant difference in mean Ct values of RNU6B, hsamiR-26b, and hsa-miR-16 between the group of patients with colonic polyps and without colonic polyps (P-value = 0.204, 95% Cl, -2.56, 0.58). This suggests that these three microRNAs are suitable candidates for normalisation control for relative RT-PCR. Analysis with all three algorithms showed that hsa-miR-16 alone had the best stability among the three microRNAs tested, while the stability improved when miR-16 and RNU6B were combined.

Conclusion

The findings in this study could be used as a guide to other research applying the relative RT-PCR technique in measuring gene expression of microRNA in human urine samples.

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P-4

A novel mutation of the *NOTCH3* Gene in Malaysian patient with Cerebral Autosomal Dominant Arteriopathy with Subcortical Infarcts and Leukoencephalopathy (CADASIL)

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Background

Mutations in the *NOTCH3* gene are responsible for cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy (CADASIL), adult-onset hereditary angiopathy leading to ischemic episodes, vascular dementia, and other neurologic deficits. Most *NOTCH3* mutations reported so far are rigidly stereotyped, involving the gain or loss of a cysteine residue in a specific *NOTCH3* epidermal growth factor (EGF)-like repeats on chromosome 19.p13.12 [1, 3]. **Methodology**

More than 100 blood samples of unrelated Malaysian patients with clinical/neuroimaging pictures suggestive of CADASIL were referred to our laboratory for molecular investigation. Genomic DNA was extracted from whole blood using Prepito DNA Blood250 Kit (PerkinElmer). Genetic diagnosis of CADASIL was performed by targeted PCR amplification of exons 3, 4, 5, 6, 11, 16 and 19, including exon-intron boundaries of the NOTCH3 gene, followed by Sanger sequencing to detect causative mutations. Sequence analysis was performed using 3500 Genetic Analyzer (Applied Biosystems). To confirm the pathogenicity of new nucleotide variants, the identified sequence variants were assessed against several public databases and *in silico* prediction tools such as VarSome in accordance with ACMG guidelines.

Results and Discussion

Out of the 108 samples tested, 23 samples were found to harbour variants suggestive of CADASIL. Using PCR-directed sequence analysis, a total of seven mutations were identified. One was a novel missense mutation, C260S and six previously reported mutations, R110C, R141C, R182C, C194S, R544C and R1006C were listed in the Human Genome Mutation Database (HGMD). In this study, a 51-year-old female patient with novel missense mutation C260S is presented with sudden onset of left-sided weakness and MRI brain scan findings suggestive of CADASIL. VarSome predicted C260S as pathogenic, located in a mutational hotspot and two different missense changes determined to be pathogenic have been seen before. The nucleotide position was strongly conserved and the variant was not found in gnomAD, a population database for normal individuals. Furthermore, the pathogenicity score was 95% from multiple *in silico* predictors. This result is consistent with the CADASIL diagnosis for this patient.

Conclusior

Careful assessments of genealogical, clinical, and neuroimaging data in patients with lacunar stroke can aid in selecting patients with a high probability of finding mutations by genetic screening [2]. In our samples, 21% of Malaysian patients with 'clinically suspected' CADA-SIL received the definitive molecularly proven diagnosis. In contrast, if clinical suspicion is high, the remaining undiagnosed patients should be recommended for further testing of NOTCH3 exons, which were not included in this study. In addition, the discovery of new mutations expands the genetic spectrum of *NOTCH3*-related diseases, which will contribute to further study of this disease in the future. Correct diagnosis is not only important for the management of the patient but also crucial for genetic counselling and early diagnosis of at-risk family members.

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P-42

Screening of Lactobacillus species in stingless bee honey

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Background

Stingless bee honey is a natural substance produced by the stingless bee of genera Meliponini. Compared to honeybee honey, stingless bee honey has a significantly higher moisture content, water activity, and ash content, while the pH and total soluble solid content are slightly lower. Other components such as proteins, amino acids, enzymes, organic acids, mineral elements, and vitamins are also present in small amounts. Being rich in nutrients, honey provides a suitable environment for microflora growth. Microflora in honey includes yeasts, moulds, and bacteria with bacteria being the major microbes associated with stingless bee honey. Many studies have proven the presence of probiotics amongst the microflora, which includes Lactobacillus, Lactococcus, and Bifidobacterium. This finding has provided additional commercial value to stingless bee honey. Hence, this study aims to screen the presence of Lactobacillus spp., one of the most accepted probiotics, in stingless bee honey of Heterotrigona itama collected from a plantation in Mantin, Negeri Sembilan.

Methodology

Samples were randomly obtained from July to November 2018 from the same plantation area. The samples were cultivated in lactobacilli de Man, Rogosa, Sharpe (MRS) broth before being subjected to isolation on MRS agar. This step was carried out 3 times to further isolate the potentially different species by differentiating them according to their colony morphology. 5 colonies isolated from each month were subjected to DNA extraction using QiaAmp DNA kits. The colonies were identified via polymerase chain reaction (PCR) using a set of primer pairs targeting the region of 16S rRNA; LF 5'-AGCAGTAGGAATCTT CCA-3' and LR 5'-ATTACACCGCTACACATG-3', targeting amplicon sizing 430bp. PCR products were sent to 1st Base Laboratories for sequencing followed by sequence alignment using BLAST on the NCBI website.

Results and Discussion

Growth on MRS agar showed the growth of a few species that could be differentiated by their colony morphologies. Colonies colour ranged from whitish to brownish yellow, while other characteristics observed were sliminess, size, and shape; where there were round and slightly oval shape. Most samples subjected to PCR gave out clear single band sizing around 400bp during gel electrophoresis, while some other gave two to three, clear but faint bands ranging around the same size; 400 -500bp. This could be explained by mixed species that co-inhibit the same honey pot. Sequencing and alignment of PCR products in the database showed that several Lactobacillus species repeatedly emerged on the list. This suggested that these species are dominant in the stingless bee colony of that plantation. Said species were Lactobacillus kefiri strain JK5, Lactobacillus timberlakei strain HV, Lactobacillus malefermentans JCM 12497, Lactobacillus kefiranofaciens strain IMAU98303 and Lactobacillus salivarius strain JCM1046.

Conclusion

This finding confirms that stingless bee honey provides a good variety of beneficial bacteria especially from *Lactobacillus* strains.

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P-43

Interaction of UV-VIS optical properties towards enterovirus 71 RNA for detection of hand foot mouth disease using optical spectroscopy

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Background

Hand-foot-mouth disease (HFMD) is a common infectious disease that occurs in children under 5 years old. The disease is transmitted through nasal fluids, saliva and contact act with surfaces contaminated with HFMD patients. HFMD is caused by viruses known as Enterovirus A71 (EV71). Generally, all viruses are biological things that contain specific genes of deoxyribonucleic acid (DNA) and ribonucleic acid (RNA). In this project, RNA extraction is used instead of DNA. RNA extraction is safer, stays inactivated at room temperature, and has resistance to ultraviolet sensitivity compared to DNA. Various types of light with different wavelengths are represented in the electromagnetic spectrum, and this research used Ultraviolet (UV) until Visible (VIS) spectra. This project aims to study the optical properties of EV71 RNA via UV-VIS spectroscopy for HFMD detection and to differentiate the UV-VIS light response between various samplings of EV71 RNA through spectra analysis. UV-VIS spectroscopy requires specialized methodologies to be performed. This project was implemented using absorption, transmission, and reflectance methodologies.

Methodology

In this study, EV71 RNA with different concentrations were used. Optical spectroscopy in the ultraviolet-visible (UV-VIS) range was used to observe the optical properties of all the RNAs, including absorbance, transmission and reflectance. The Ocean View software was used to capture the spectra of all honey. The spectra were collected more than 10 times for each concentration to improve the accuracy of the results. The average of the results was plotted, and analysis was performed for all spectra results.

Results and Discussion

As a result of this project, the value of absorbance and reflectance is directly proportional to the concentration whereas the value of transmittance is inversely proportional to the concentration. The results of this experiment were supported by theoretical and previous studies. The data of this research need continue to be discussed and improved from time in developing a diagnostic instrument that is responsive, mobile, lowcost, and accurate. A rapid, adaptive test, such as a "test kit" capable of identifying the EV71, will significantly improve the detection of HFMD. Thus, the annual number of HFMD cases can be controlled since the appropriate treatment can be administered due to the early detection of HFMD in the early stages.

Conclusion

Three optical properties, which are absorbance, transmission, and reflectance were studied for all EV71 RNA using optical spectroscopy in the UV-VIS ranges. An analysis was performed on the spectra results to compare the mean value of the peak spectra. Absorption and reflection have demonstrated a similar relationship between the sample concentration used and the absorbance or reflectance values. This means, when the sample concentration was high, the absorbance or reflectance value will also be increased. Contrarily, the association between sample concentration and transmittance were different. This signifies that the transmission value was low when the sample concentration was high.

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P-44

Analysis of microsatellite instability in colorectal cancer patients from Hospital USM

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Background

Microsatellites are short, repeating DNA segments vulnerable to mismatches during replication. Mismatch repair (MMR) protein complexes often correct DNA mismatch errors. DNA errors build-up due to the MMR system's loss of function, causing microsatellite instability (MSI), a genetic hypermutability disorder. MSI is a molecular phenotype due to a defective DNA MMR system. Recent years have seen a rise in interest in DNA MMR deficiency as a critical biomarker to predict the therapeutic efficacy of immune checkpoint inhibitors for various malignant neoplasms. The study aims to investigate the profile of MSI status in colorectal cancer (CRC) patients from Hospital USM.

Methodology

The formalin-fixed, paraffin-embedded tissue blocks from archival colon cancer samples and adjacent normal tissue (n = 43) were randomly selected between 2018 and 2020. Genomic DNA extracted from tumors and corresponding normal tissues were used for MSI analysis using the Promega panel (5 mononucleotide markers: BAT-25, BAT-26, NR-21, NR-24, and MONO-27; and 2 pentanucleotide markers: Penta C and Penta D).

Results and Discussion

Of the 43 tissue samples studied, 21 samples came from men and 22 came from women. The median age of patients in the study group was 62 years. Four tumors (9.3%) demonstrated high MSI (MSI-H), one demonstrated low MSI (2.3%), and the remaining 38 (88.4%) tumors were microsatellite stable (MSS). Our finding was comparable with other studies which show that the incidence of MSI in tumors in CRC varies from 8.8% to 20.3%. Interestingly, patients in the MSI-H group were younger, with a median age of 39 years old compared with the study population with a median age of 62 years old. In constrast, a median age of 63 years was observed in the MSS group. According to a study, patients with CRC who had MSI-H were diagnosed earlier than those with MSS (1). Most selected tissue samples arise from the distal colon whereas smaller percentage were from proximal colon (16%). Notably, the incidence of proximal lesions in the MSI-H group (75%, 3/4) was higher than in the MSS group (9%, 4/39). The findings of researcher from several countries have demonstrated that MSI is more frequent in patients with tumors localization in the proximal of the colon (2), which is also in agreement with our findings. It has been reported that MSI-H tumors respond well to immunotherapy, PD-L1 (programed cell death ligand 1), with FDA approved to treat MSI-H/MMR patients.

Conclusion

This is an exploratory study employing a Promega panel to look at the MSI profile in our CRC patients. To associate clinical characteristics with MSI status, it is necessary to do further research with larger sample size. It would be beneficial to screen patients with MSI-H status which has a potential role in treatment management.

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P-46

Antimicrobial activity of astaxanthin extracts of wild type and mutant Xanthophyllomyces dendrorhous

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Background

There is worldwide concern about the increase in antimicrobial resistance which affects both developed and developing countries. It is a public health issue with immense societal and economic implications [1]. Antibiotic-resistant pathogens pose an enormous threat to the treatment of a wide range of serious infections. A periodic replacement of the existing antibiotic is necessary to prevent this exponential emergence. *X. dendrorhous* extracts were signified on the benefits of astaxanthin in antioxidant activity, but less emphasized for its antimicrobial activity. This study aims to assess the antimicrobial ability of astaxanthin extract of wild type and mutant *X. dendrorhous* obtained in our research.

Methodology

Antimicrobial activity of astaxanthin extracts was evaluated through the zone of inhibition using paper disc assay against seven microorganisms of industrial importance Gram-negative bacteria: *Shigella boydii* and *Escherichia coli*, Gram positive bacteria: *Bacillus subtilis* and *Staphylococcus aureus*, yeasts: *Saccharomyces cerevisiae* and *Candida albicans*, and fungus: *Aspergillus niger*. The wild type and mutant astaxanthin extracts were prepared at a concentration of 30ug/mL and dotted on autoclaved paper discs. The experiment also included a positive control of 30ug/mL of chloramphenicol for antibacterial drugs and Miconazole as a standard antifungal drug. All the plates containing bacteria were incubated at 37°C for 24 h and that of yeast and fungi at 28°C for 48 h.

Results and Discussion

The standard antibiotic chloramphenicol produced inhibition zones of 12, 14, 9 and 8mm against *B. subtilis, S. aureus, S. boydii* and *E. coli* respectively. Meanwhile positive control Miconazole produced inhibition zones of 28, 25 and 23 mm against *S. cerevisiae, C. albicans* and *A. niger* respectively. The antimicrobial activity of both wild type and mutant extracts showed significant growth inhibition against pathogenic microbial strains. The mutant astaxanthin extract showed a slightly better inhibition effect on all the pathogens than the wild type extract, but the difference was not significant. The extracts exhibited an excellent inhibitory effect against Gram-positive *B. subtilis*

and *S. aureus*, nearly as good as positive control. They exerted a milder antimicrobial capability against Gram negative *S. boydii* and *E. coli*. Compared to the positive control, the extracts showed moderate inhibition in *S. cerevisiae*, *C. albicans* and *A. niger* for yeast and fungus. Limited antimicrobial reports on astaxanthin are mainly extracted from sources like microalga *H. pluvialis* and crustaceans. Some contradictions can be found in the literature due to the different strains used in the determination of antimicrobial activity, extraction methods, extraction solvents and extract concentration used in the assays [2].

Conclusion

Based on controls, both the wild type and mutant astaxanthin extracts showed impressive inhibition on Gram positive and Gram negative bacteria compared to yeast and fungus. The twofold use of astaxanthin extracts as antioxidants and antimicrobials has good prospects in response to the consumer demand for more sustainable and natural ingredients and products.

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P-47

The potential of Xanthophyllomyces dendrorhous wild type and mutant strains in producing UV absorbing compounds

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Background

Solar ultraviolet radiation is the culprit for molecular and genetic changes that occur to the skin. It generates reactive oxygen species and free radicals that cause skin oxidative damage leading to skin cancers, photosensitivity, and photo-aging problems. These contribute to synthetic sunscreens formulated with UVR filters that mostly show side effects on human health and ecology. Hence, research investigations were initiated upon natural resources that can produce UV-absorbing compounds known as mycosporine-like amino acids and mycosporine-glutaminol-glucoside. These molecules are commonly used by organisms growing at high altitudes to protect themselves from extreme UV exposure [1]. In this study, a basidiomycetous yeast, *Xanthophllomyces dendrorhous* wild type and its mutant strains were analyzed for their ability to produce MAA/MGGs.

Methodology

X. dendrorhous wild type and mutant strains were revived and maintained on yeast malt agar at 4°C. Synthesis of mycosporines was induced by transferring young cultures (24hr) to YM agar plates and incubated for 4 days at 22°C in an incubator (LabCompanion SIF-6000R) with a 12:12 light: dark photoperiod. For light conditions, culture plates were left exposed to PAR+UVR while for dark conditions, plates were covered with aluminium foil. After exposure, the colonies were harvested and transferred to distilled water, centrifuged, and conserved at -20° until mycosporines extraction. Samples were extracted with 10ml of DMSO solution (2%) for 24hrs at 4°C. After 24hrs, glass beads were added to samples and vigorously agitated in

a vortex mixer. Later, they were incubated for 24hrs at 45°C in a water bath and centrifuged using a microcentrifuge. The resulting supernatant was immediately measured spectrophotometrically (UV-visible) at 325 nm for mycosporines quantification. Experiments were performed in triplicate.

Results and Discussion

With the aid of a UV spectrum, the mycosporine production rate was found to be the highest at 325 nm for all four strains leading to a fixed wavelength for further measurements. Yeast colonies exposed to PAR+ UVR showed higher absorbance rates concluding more mycosporines production than those grown in a dark environment. It showed that strains frequently exposed to high UV radiation have an increased tendency of mycosporine production than those with lower intensity of UV exposure. By comparing the wild type and 3 mutant strains, the results indicated that the red, white and yellow mutant colonies initially showed almost equal mycosporines production rate as the wild strain. The production rate will be further improved by introducing different inducers, growth media and culture conditions.

Conclusion

All the four strains tested showed the detection of MAA/MGGs in the DMSO extracts. It is concluded that after photostimulation, both *X. dendrorhous* wild type and mutant strains have the ability to synthesize UV-absorbing compounds and further investigation should be done to discover its potential as a natural molecular sunscreen.

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P-50

Comparative evaluation of three methods for protein extraction efficiency from formalin-fixed paraffin-embedded tissues for proteomic analysis

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Background

Formalin fixation and paraffin embedding of tissues (FFPE) have become the most available biopsies in the hospital setting due to their ability to be stored for long periods and may be kept at room temperature without concern of deterioration or decay. However, this fixation process causes a protein alteration known as "cross-links." Consequently, it made it challenging to extract proteins and characterize individual proteins. The study of proteomics is growing quickly because it has the potential to enhance disease diagnosis, risk assessment, prognosis, and therapy targeting. Consequently, as a researcher, it is important to investigate strong and efficient protein identification approaches and improve all aspects of protein analysis, such as the protein extraction method from FFPE tissue and protein digestion for mass spectrometry.

Methodology

For this research, FFPE human tonsillitis blocks were chosen for this research to compare the efficiency of three different protein extraction methods, including sonication, RapiGest, and Sodium

Deoxycholate, from Formalin Fixed Paraffin Embedded (FFPE) tissue for proteomic analysis. These sections were cut into 4 μ m thin slices of tissue to facilitate the deparaffinization process. Three sets of unstained FFPE tonsil tissue sections were excised into 1.0 cm2 for the triplicate results. For the quantification analysis, Nanodrop 2000 was used to measure the protein concentration from the extracted protein. While the determination of the protein quality, SDS-PAGE analysis was done with three different protein extraction methods. The proteins of interest are digested with an enzyme, trypsin, and the resultant peptides are examined using mass spectrometry. Protein identification will be accomplished using LC-MS/MS analysis of protein extracted from three sets of each extraction technique obtained from FFPE tonsillitis samples.

Results and Discussion

The protein concentration of the samples was determined using the nanodrop 2000c. From the sonication method, the protein concentration was extremely low and inconsistent. The sodium deoxycholate method produced a greater protein concentration than RapiGest. Next, the protein quality was evaluated using SDS-PAGE following the application of the three protein extraction methods for sample fragmentation. The intensity of the band for RapiGest and Sodium Deoxycholate can be observed at 1 cm². However, a very faint band was observed for the sonication method. This shows that the highest abundance of protein loaded into the gel is from the Rapi-Gest and Sodium Deoxycholate, while the lowest is from the sonication method. All the proteins were then analysed via LC/MS and identified using Peak 7.5 software. A total of 668 proteins were successfully identified using the sodium deoxycholate method. In addition, a total of 387 protein samples were obtained using the sonication method. However, for the RapiGest method, the number of proteins acquired from the samples was only 89, which was the lowest. Based on the results obtained from the software, 14 proteins were found to overlap in all three protein extraction methods.

Conclusion

In conclusion, the most efficient protein extraction method is sodium deoxycholate because the protein concentration is the highest. Adding to that, the number of proteins identified using this method is also the highest.

P-51

Optimization of expression and purification of soluble *E. bieneusi* B7XHF2 recombinant protein in *E.coli*

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Background

Worldwide reports of human infections with *Microsporidia* mainly include immunocompromised patients [1]. One of the most prevalent microsporidian species to infect HIV/AIDS patients is *Enterocytozoon bieneusi* [2]. Although stool testing is the most reliable method to diagnose microsporidiosis, the infection is underdiagnosed in patients with disseminated microsporidiosis due to the absence of microsporidia spores in their stool samples. Our previous study has identified two circulating antigens of diagnostic value from the serum of immunocompromised patients. In this study, we aimed to express and purify *E.bieneusi* recombinant protein B7XHF2 in *Escherichia coli* as a potential diagnostic biomarker for the detection of disseminated microsporidiosis.

Methodology

The B7XHF2 gene was cloned in the pET-32a(+) vector, and the recombinant plasmid was transformed into C41(DE3) *E.coli* host cell.

Small scale expression in 200ml Terrific broth (TB) was performed to optimize the parameters of protein expression namely IPTG concentration and post-induction incubation period. Subsequently, protein expression was carried out in 2L TB medium, at a temperature of 37°C, 200 rpm before the temperature was lowered to 28°C for 4 hours post IPTG induction. The harvested cells were pelleted and lysed by sonication in a lysis buffer containing protease inhibitors and lysozyme. The filtered protein lysate was then subjected to affinity purification using Ni-NTA resin at a ratio of 1:10 (resin:lysate). Contaminating proteins were removed by gradient washing using washing buffer containing 20mM, 30mM and 40mM imidazole concentration, and finally the targeted protein was eluted by adding elution buffer with 250mM imidazole. Eluted fractions were pooled and concentrated to 300ul and the protein concentration was measured by the RCDC method (BioradTM). Anti-histidine western blot was carried out to verify the presence of the targeted protein.

Results and Discussion

In this study, the rB7XHF2 protein was highly soluble when induced with 0.5mM IPTG compared to 1.0mM IPTG. There was no significant difference in terms of protein expression when incubation was performed at 2,3,4 and 16 hours post-induction, therefore 4 hours was chosen as the best incubation period. The appearance of a 41-kDa protein on anti-histidine western blot confirmed the expression of rB7XHF2 protein, which was close to the theoretical size of 41.7 -kDa.

This preliminary study has successfully expressed the B7XHF2 recombinant protein in *E.coli* C41(DE3) host cells. Further work is needed to improve the yield as well as purity of the target protein for diagnostic applications.

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P-54

In silico selection of aptamers to cobra venom cytotoxin

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Background

Snakebite envenomation has been listed as one of the neglected tropical diseases (NTDs) by the World Health Organization (WHO) since 2017 [1]. Cytotoxin (CTX), one of the most abundant venom components, was deemed responsible for the envenomation's local dermonecrosis. Several studies have indicated that the CTX induces cell death through different pathways, yet the molecular mechanism is inconclusive [2]. There is also limited treatment against dermone-crosis. Therefore, there is a need to search for biotherapeutics against CTX-induced dermonecrosis. In this study, we exploit the potential of tRNA-based aptamers as alternate toxin-neutralizing molecules using in silico approach.

Methodology

GtRNAdb database was used to identify the CTX-binding tRNA-based aptamers for the possible interacting regions with CTX [3]. The sequences of the aptamers were subjected to phylogenetic tree analysis to categorize the tRNAs into different clusters using the Clustal Omega. The tertiary structures of the consensus aptamer sequences were modeled using the RNA Composer. Later, the molecular docking of these aptamers and CTX was performed using HADDOCK web server. In addition, RING analysis was performed on each aptamer-

CTX complex to determine the interacting amino acids present on the CTX and the type of interactions.

Results and Discussion

Altogether, our results suggested six clusters of consensus aptamer sequences from the phylogenetic tree analysis. Among the six aptamer models generated from their respective consensus sequences, three aptamers demonstrated significant binding to CTX. Additionally, the RING analysis revealed that there were four common interacting regions between the six CTX-aptamer complexes. These four interacting regions were consistent with the CTX's epitopes in our unpublished work. Moreover, the high binding affinity between the aptamer-CTX complexes also suggested that they were potential aptamers to antagonize CTX's action.

Conclusion

The current work provides insights into *in silico* approach to design aptamers that antagonize CTX-induced dermonecrosis. Three tRNA-based aptamers have been identified as potential aptamers, which showed strong interaction with CTX's epitopes.

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P-55

IgG2a subclass response of mice towards vaccination of Strongyloides stercoralis antigens

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Background

It has been tried and proven that vaccination is still the best strategy to combat infectious diseases. However, till date, there are still no vaccines against strongyloidiasis, caused by *Strongyloides stercoralis*. Approximately 100 million people are estimated to be infected worldwide, most commonly in tropical and subtropical regions. There are very few studies on vaccine candidates for strongyloidiasis. To date, there is only one study that uses recombinant protein from *S. stercoralis*, Ss-IR [1]. Furthermore, another study performed immunoscreenings with anti-human IgE antibody of an *S. stercoralis* complementary DNA (cDNA) library, with the results that the cDNA clone A-133 having the highest diagnostic potential [2]. Seeing the protein Ss-IR and A-133 having high potential as vaccine candidate against

strongyloidiasis, this study aims to investigate the effects of vaccination with rSs-IR and rA-133 on levels of serum IgG2a antibody. **Methodology**

rSs-IR and rA-133 conjugated with histidine tags will be expressed using *Escherichia coli* C41 (DE3) and purified via Ni-NTA affinity column chromatography. The purified antigens will be used for immunization of female Balb/c mice that will be divided into four test groups, namely unvaccinated control (n=6), adjuvant-only control (n=8), rA-133 treatment (n=13) and rSs-IR treatment (n=12). Priming dose will be administered via intraperitoneal route adjuvanted with Complete Freund's Adjuvant (CFA), whereas subsequent booster doses will be administered via subcutaneous route adjuvanted with Incomplete Freund's Adjuvant (IFA). Serum samples will be obtained via blood bled via tail snip. After that, ELISA will be performed on serum samples to monitor the level of subclass IgG2a antibodies. Results and Discussion

The IgG2a levels in mice sera increased with each administration of rSs-IR and rA-133 respectively. Furthermore, rA-133 elicited a greater increase in IgG2a serum levels compared to rSs-IR. IgG antibodies are primarily involved in anti-pathogenic adaptive immune response and long-term protection, and IgG isotype antibody production dictates the robustness of immune response following antigen immunization. The finding of a higher IgG2a production stimulated by rA-133 is a favorable outcome for the antigen as vaccine candidate, due to the important roles IgG2a play in other parasitic diseases. For instance, it has been shown that IgG2a can activate macrophages that traps tissue-migrating helminths larvae thus immobilizing it [3]. **Conclusion**

In conclusion, this profile of high IgG2a expression shown by rA133 in this study is an important finding since more studies into parasitic carbohydrate-based vaccines are being conducted in order to generate similar antibody profiles. Furthermore, the result of this study provides a rough clue in figuring out the immune responses towards a steronalis

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