Chromatographic Analyses, Virtual Screening and Pharmacokinetics of Yellow Malaysian Rambutan (Nephelium lappaceum L.) Fruit Epicarp Extracts Reveal Potential Antibacterial Compounds

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Abstract

The emergence and spread of antimicrobial resistance have been of serious concern on human health and the management of bacterial infectious diseases. Effective treatment of these diseases requires the development of novel therapeutics, preferably free of side effects. In this regard, natural products are frequently conceived to be potential alternative sources for novel antibacterial compounds. Herein, we have evaluated the antibacterial activity of the epicarp extracts of the Malaysian cultivar of yellow rambutan fruit (*Nephelium lappaceum L.*) against six pathogens namely, *Bacillus subtilis*, methicillin-resistant *Staphylococcus aureus* (MRSA), *Streptococcus pyogenes*, *Pseudomonas aeruginosa*, *Klebsiella pneumoniae* and *Salmonella enterica*. Among a series of solvent extracts, fractions of ethyl acetate and acetone have revealed significant activity towards all tested strains. Chemical profiling of these fractions, via HPLC, LC-MS and GC-MS, has generated a library of potential bioactive compounds. Downstream virtual screening, pharmacological prediction, and receptor-ligand molecular dynamics simulation have eventually unveiled novel potential antibacterial compounds, which can be extracted for medicinal use. To this end, we report novel mechanistic aspects of these compounds in competitively inhibiting the ATP-binding domain of the DnaK chaperone of *P. aeruginosa* and MRSA. Our work takes a step forward to discover antimicrobials capable of perforating the barrier of resistance posed by both the gram positives and the negatives.

Introduction

Antimicrobial resistance (AMR) has been projected as one of the serious global concerns with issues in the management of infectious diseases caused by Multidrug Resistant (MDR) bacterial pathogens\(^1\). These pathogens have been reported to cause 700,000 deaths each year which is estimated to cross over 10 million by 2050\(^2\). Such alarming rise can be attributed mostly to the prevalent misappropriation of antibiotics in human healthcare systems\(^3\) delineated by the overuse and poor appropriate prescriptions coupled with the lack of new drug development, which have reduced the efficiency of antibiotics as the resistant strains rapidly increases\(^4\). This necessitates the urgency for unearthing alternate natural therapeutics focused on natural products to be exploited as repertoires of natural bioactive compounds, preferably devoid of side effects and hence, potential for future drug development.

Natural products, frequently discovered with potent antimicrobial potential, have been used as alternatives with the hope of eliminating the use of synthetic antibiotics, since a long time\(^5\). Besides serving a widespread range of secondary plant metabolites, for instance, alkaloids, tannins, flavonoids and phenolic compounds, with remarkable antimicrobial effects\(^5\), many plant products have been reported for their antimicrobial potentials, namely, citrus peel\(^7\), grape seed\(^8\), cranberry pomace\(^9\), pomegranate peel\(^10\) and passion fruit seed\(^11\). In this regard, the fruit rambutan (*Nephelium lappaceum L.*) has gained attention due to its vast range of bioactive constituents including, but not limited to, vitamin C, vitamin E, carotenes, xanthophylls, tannins and phenolic compounds like, geraniin, ellagic acid, quercetin, corilagin and rutin\(^12\).
While the broad range of biological activities of rambutan include anticancer, antiproliferative and anti hypercholesterolemic properties\textsuperscript{13,14}, few researchers have even reported its \textit{in vitro} antibacterial activities. For instance, Malini & Maheshkumar\textsuperscript{15} have disclosed significant antibacterial activity of rambutan fruit sap extracts towards \textit{Pseudomonas aeruginosa} while Bhat and Al-daihan\textsuperscript{16} revealed antibacterial activities of its seeds extracts against \textit{Staphylococcus aureus}, \textit{Streptococcus pyogenes}, \textit{Bacillus subtilis}, \textit{Escherichia coli} and \textit{P. aeruginosa}. Moreover, antibacterial potential of rambutan peel extracts have also been reported against \textit{Vibrio cholerae}, \textit{Enterococcus faecalis}, \textit{S. aureus} and \textit{Staphylococcus epidermidis}\textsuperscript{17}. Furthermore, Sekar \textit{et al.}\textsuperscript{18} comparatively evaluated the antibacterial potency of red and yellow rambutan fruit peels against \textit{S. aureus} and \textit{S. pyogenes}, to reveal better efficacy of the latter extracts against the tested pathogens.

In this study, we have delineated a stepwise approach of determining the efficacy of the crude extracts of the epicarp of yellow Malaysian rambutan against clinically important MDR bacterial pathogens e.g. \textit{B. subtilis}, methicillin-resistant \textit{S. aureus} (MRSA), \textit{S. pyogenes}, \textit{P. aeruginosa}, \textit{K. pneumoniae} and \textit{S. enterica}. Further chemical profiling of the extracts through HPLC, LC-MS and GC-MS revealed their potential chemical determinants which were screened virtually to pharmacologically unveil novel bioactive compounds through molecular dynamics simulation. To this end, a competitive inhibition of the ATP-binding domain of the DnaK chaperone of \textit{P. aeruginosa} and MRSA, by our shortlisted compounds, portrayed novel antibacterial mechanism, representative of targeting both the gram-positive and the -negative pathogens.

**Results**

**Variable antibacterial activity of crude extracts by disc diffusion**

A preliminary screening for the antibacterial activities of the yellow-variety Malaysian Rambutan epicarp crude extracts was assessed through disc diffusion assay. Only freshly prepared solutions were used for all the tested pathogens (TP) for the crude extracts. Throughout such trials, the solvent control (SC), DMSO, showing no inhibition zone, did not exercise antibacterial activity against the TP. Moreover, none of the sequential extracts exhibited activity against the TP (Table 1, Fig. S1), while in case of direct extracts, the picture was quite different. Extracts of ethyl acetate (EA) displayed visible activity against MRSA, \textit{B. subtilis} and \textit{S. enterica} but did not show activity against the rest of the TP (RTP). Again, acetone (AC) extracts exhibited markedly prominent activity against \textit{B. subtilis} along with visible activity against \textit{S. pyogenes} and \textit{P. aeruginosa}, which was not observed among the RTP (Table 1, Fig. S2).
Table 1
Antibacterial activity of the sequential and direct crude extracts of *N. lappaceum* by disc diffusion

| Exports | B. subtilis | MRSA | S. pyogenes | P. aeruginosa | K. pneumonia | S. enterica | S.C | P.C |
|---------|-------------|------|-------------|---------------|--------------|-------------|-----|-----|
| EA(S)   | -           | -    | -           | -             | -            | -           | -   | ++  |
| AC(S)   | -           | -    | -           | -             | -            | -           | -   | ++  |
| EA(D)   | +           | +    | -           | -             | -            | +           | -   | ++  |
| AC(D)   | ++          | -    | +           | +             | -            | -           | -   | ++  |

+: Visible; ++: Prominent; -: No activity; EA: Ethyl acetate; AC: Acetone S: Sequential; D: Direct; S.C: Solvent control (DMSO); P.C: Positive control (Gentamicin 10 µg)

Qualitative antibacterial screening of crude extracts via Broth dilution

Broth dilution method was used for evaluating the antibacterial potential of yellow fruit epicarp crude extracts and calculating the viability percentage of every TP (Table 2). This in turn illustrated the percentages of antibacterial potential of EA and AC fractions from the sequential & direct extracts, against the TP, at concentration of 250 µg/mL. For the sequential extracts, the fraction from AC exhibited the highest percentage (90) of antibacterial activity against *P. aeruginosa* followed by 71% against MRSA while that from EA exhibited 59% effect against *S. pyogenes*. No significant results were observed either against the RTP (Fig. S3 a-f) or for the remaining solvent extracts namely, chloroform (CF), ethanol (ET), methanol (MT) and aqueous (water, WT) extracts against all TP (*Table S2*). In the case of direct extracts with the six solvents used, fractions of EA exhibited inhibition of 80% against *B. subtilis* along with 60, 62, 73 and 72% for MRSA, *P. aeruginosa, S. enterica* and *K. pneumonia*, respectively, without any positive results against *S. pyogenes*. Notably, all tested pathogens were inhibited by AC fractions and the percentage of antibacterial effects were 75, 90, 70, 70, 60 and 75, respectively for MRSA, *B. subtilis, S. pyogenes, P. aeruginosa, S. enterica* and *K. pneumonia* (*Fig. S3 g-l*).
Table 2  
Screening of antibacterial effect of sequential and direct extracts by broth dilution method

| Microorganisms | EA(S) | EA(D) | AC(S) | AC(D) | S.C | P.C |
|----------------|-------|-------|-------|-------|-----|-----|
| B. subtilis    | -     | 20    | -     | 10    | 98  | 0   |
| MRSA           | -     | 40    | 29    | 25    | 98  | 0   |
| S. pyogenes    | 41    | -     | -     | 30    | 98  | 0   |
| P. aeruginosa  | -     | 38    | 10    | 30    | 98  | 0   |
| K. pneumonia   | -     | 28    | -     | 25    | 98  | 0   |
| S. enterica    | -     | 27    | -     | 40    | 98  | 0   |

EA(S): ethyl acetate sequential; AC(S): acetone sequential; EA(D): ethyl acetate direct; AC(D): acetone direct; S.C: solvent control: DMSO (< 1%); P.C: positive control (Gentamicin 10 µg); -: Low activity (less than 50%).

The results of EA and AC fractions portrayed notable antibacterial efficiency and were, thus, processed for identification of bioactive compounds via HPLC, LC-MS and GC-MS analyses.

Revelation of antioxidants from crude extracts using HPLC-UV

For a preliminary identification of the basic antioxidants present in the Malaysian yellow-rambutan epicarp extracts, HPLC-UV was conducted. All compounds with known antioxidant capacities were identified in comparison with standard phenolic compounds. The identified compounds and their quantification, along with their specific peak position and retention time (Rt) in chromatogram, are shown in Table 3 and Fig. S4. Only three compounds were identified in the EA extract namely, malic acid, vitamin C and chlorogenic acid along with three more in the AC extract. These are epigallocatechin gallate, catechin hydrate and quercetin.
Table 3
Identified compounds in *N. lappaceum* ethyl acetate and acetone sequential fractions using HPLC-UV.

| Sample Extract | RT (min) | Identified compounds       | Peak Area  |
|----------------|----------|---------------------------|------------|
| EA (S)         | 12.9     | Malic acid                | 40.441     |
|                | 4.8      | Vitamin C                 | 1717.201   |
|                | 6.2      | Chlorogenic acid          | 190.156    |
| AC (S)         | 2.9      | Malic acid                | 4.938      |
|                | 4.8      | Vitamin C                 | 1226.316   |
|                | 6.2      | Chlorogenic acid          | 134.587    |
|                | 8.8      | Epigallocatechin gallate  | 205.717    |
|                | 10.7     | Quercetin                 | 85.112     |
|                | 20.7     | Catechin Hydrate          | 44.169     |

EA (S): Ethyl acetate sequential fraction; AC (S): Acetone sequential fraction

Exploration of other chemical determinants through Liquid Chromatography–Mass Spectrometry (LC-MS) Analysis

For a fast, mass-directed exploration of the compounds possibly present in the rambutan epicarp, its EA and AC sequential crude extracts were subjected to LC-MS analysis (Fig. S5 and Fig. S6) which revealed the presence of 54 and 44 compounds, respectively (Table S3 & S4). They were matched with the identity of known molecules on the Metlin database, keeping a threshold Molecular Formula Generator (MFG) scores above 86% along with a ± 2% difference. The compounds above the mentioned cut-off were further explored for their biological activities and carried forward for virtual screening. Notably, 31 compounds from both sets of EA and AC extracts have not been reported till date with any antibacterial activities (Table S3 & S4).

Identification of volatile constituents by Gas chromatography–mass spectrometry (GC-MS)

To identify any volatile organic compounds, present in the EA and AC extracts of Malaysian yellow-variety *N. lappaceum* epicarp, they were exposed to GC-MS analysis (Fig. S7 a-d). Most of the compounds from EA and AC fractions, extracted directly, have been reported with antibacterial activities (Table 4). On the contrary, the compounds of these fractions from sequential extraction have not been reported for any such activity. The chromatogram of these compounds showed mentionable area % scores (above 0.5%) for 3-Methyl-1,2-diazirine (compound 1) and Card-20(22)-enolide, 3-{[(6-deoxy-3,4-O-methylenehexopyranos-2-ulos-l-yl) oxy]-5,11/14-trihydroxy-12 -oxo-, (3-beta, 5-alpha, 11-alpha) (compound 2) in the EA extract while, the AC extract showed the presence of Silane, [(3alpha,5beta,20S)-
pregn-11-ene-3,11,17,20-tetrayl tetrakis(oxy)] tetrakis [trimethyl] and 2,2-Bis[4-{(4,6-dichloro-1,3,5-triazin-2-yl) oxy] phenyl]-1,1,3,3,3-hexafluoropropane (Table 4). Of these, compound 2 is known as Eplerenone (Fig. 1) and was found to be an important one in the upcoming analyses.
Table 4
Compounds existing in *N. lappaceum* ethyl acetate and acetone (sequential & direct) extract identified by GC-MS analysis

| No | Extracts | Identified compounds                                                                 | Molecular Formula | R.T. (Min) | Area % | Antibacterial activity report |
|----|----------|---------------------------------------------------------------------------------------|-------------------|------------|--------|------------------------------|
| 1  | EA(S)    | 3-Methyl-1,2-diazirine                                                               | C₂H₄N₂           | 3.02       | 0.776  | Not reported                 |
| 2  |          | Card-20(22)-enolide, 3-{(6-deoxy-3,4-O-methylenehexopyranos-2-ulos-1-yl)oxy}-5,11/14-trihydroxy-12-oxo-,(3-beta.,5-alpha.,11-alpha.) | C₃₀H₄₀O₁₁        | 3.00       | 3.073  | Not reported                 |
| 3  | AC(S)    | Silane, [[(3alpha,5beta,20S)-pregn-11-ene-3,11,17,20-tetrayl] tetrakis(oxy)] tetrakis [trimethyl] | C₃₃H₆₆O₄Si₄     | 41.85      | 5.257  | Not reported                 |
| 4  |          | 2,2-Bis[4-[(4,6-dichloro-1,3,5-triazin-2-yl)oxy]phenyl]-1,1,3,3,3-hexafluoropropane | C₂₁H₈Cl₄F₆N₆O₂  | 3.00       | 4.241  | Not reported                 |
| 1  | EA(D)    | Phenol, 2,4-bis(1,1-dimethylethyl)                                                   | C₁₄H₂₂O          | 10.74      | 38.698 | 19                           |
| 2  |          | Curlone                                                                               | C₁₅H₂₂O          | 14.44      | 4.233  | 20                           |
| 3  |          | Ar-tumerone                                                                           | C₁₅H₂₀O          | 18.65      | 0.857  | 21                           |
| 4  |          | Stigmasterol                                                                          | C₂₉H₄₈O          | 17.87      | 1.66   | 22                           |
| 5  |          | n-Hexadecanoic acid                                                                  | C₁₆H₃₂O₂         | 48.78      | 3.2    | 23                           |
| 6  |          | 3,7,11-Tridecatrienenitrile, 4,8,12-trimethyl                                         | C₁₆H₂₅N          | 23.94      | 2.567  | Not reported                 |
| 7  |          | 2-Methoxy-1,3-dioxolane                                                               | C₄H₈O₃           | 39.4       | 0.563  | Not reported                 |
| 8  |          | 5H-Cyclopropa (3,4) benz(1,2-e) azulen-5-one, 1,1a-à,1b-à,4,4a,7a-à,7b,8,9,9a-decahydro-7b-à,9-à,9a-à-trihydroxy-3-hydroxymethyl-1,1,6,8-à-tetramethyl-4a-methoxy-, 9,9a-didecanoate | C₄₁H₆₆O₈         | 58.69      | 2.34   | Not reported                 |

EA(S): Ethyl acetate sequential; AC(S): Acetone Sequential; EA(D): Ethyl acetate direct; AC(D): Acetone direct
| No | Extracts | Identified compounds                              | Molecular Formula | R.T. (Min) | Area % | Antibacterial activity report |
|----|----------|--------------------------------------------------|-------------------|------------|--------|-------------------------------|
| 1  | AC(D)    | Phenol, 2-methoxy-3-(2-propenyl)                 | C$_{10}$H$_{12}$O$_2$ | 10.73      | 0.678  | Not reported                  |
| 2  |          | Phenol, 2,4-bis(1,1-dimethylethyl)               | C$_{14}$H$_{22}$O  | 14.44      | 0.912  | 19                            |
| 3  |          | Alpha-Tocopherol                                 | C$_{31}$H$_{52}$O$_3$ | 44.65      | 1.612  | 24                            |
| 4  |          | Curlone                                          | C$_{15}$H$_{22}$O  | 18.65      | 1.156  | 20                            |
| 5  |          | 1,3-Dioxolane, 2-pentadecyl                      | C$_{21}$H$_{40}$O$_4$ | 57.36      | 3.06   | Not reported                  |
| 6  |          | Ar-tumerone                                      | C$_{15}$H$_{20}$O  | 17.87      | 4.081  | 21                            |

EA(S): Ethyl acetate sequential; AC(S): Acetone Sequential; EA(D): Ethyl acetate direct; AC(D): Acetone direct

**Short listing of antibacterial compounds via virtual screening and pharmacokinetics**

Ensuing virtual screening of 91 chemical compounds obtained through the chromatographic analyses, 41 of them, having binding energy less than −7 kcal/mol, were considered as good binders (Table S9; Fig. 1, 2A). Among these, potential drug candidates were chosen based on the predicted pharmacokinetic properties. For example, good gastrointestinal (GI) absorption, bad BBB permeability, and non-P-glycoprotein (PGP) substrates were preferred for absorption properties. For metabolism, non-cytochrome P450 inhibitors were preferred. Violations on drug-likeness rules were, of course, not favored. To cater to the need, five rules have been considered, namely, the Lipinski, Ghose, Veber, Egan, and Muegge rules$^{25-29}$. Lastly, higher bioavailability scores were preferred. The Abbot Bioavailability Score utilized herein was to predict chances of drug bioavailability to be more than 10% upon oral intake$^{30}$.

Among the virtually screened compounds, catechin (C), eplerenone (E) and oritin-4-beta-ol (O) stood out to be good binders with their average binding energies being −8.205, -7.980 and −7.190 kcal/mol, respectively for *S. aureus* (*Sa*) and *P. aeruginosa* (*Pa*) DnaK proteins. C, E, O also exhibited good predicted pharmacological properties except that C is a PGP substrate (Fig. 1, Table S9). The binding conformations of C (Fig. 2B), E (Fig. 2C), and O (Fig. 2D) to both DnaK proteins of *Sa* (*SaD*) and *Pa* (*PaD*) showed potential structural competitive inhibition of ATP binding at the docking pocket. Moreover, rich electrostatic interactions (Fig. 2E) were observed in C and O, but not in E, having only one intermolecular hydrogen bond.

**Validation of inhibitory effects of selected compounds by Molecular Dynamics Simulation**
Molecular Dynamics (MD) simulations for 10 nanoseconds were carried out for C, E, and O Ligand-SaD/PaD complexes to observe ligand-receptor interactions. Over the course of MD simulations, the ligands were retained in the docking pocket of respective DnaK receptors, except for C in SaD system (CSaD) of which the ligand seemed to be escaping from the initial binding pocket (Fig. 3A; B). Moreover, the upper part of the DnaK NBD domain was completely disintegrated in CSaD. Besides, the total number of receptor-ligand intermolecular hydrogen bonds were maintained stably at around 4 and 5 in *P. aeruginosa* DnaK complexed with C (CPaD) and O (OPaD) respectively, and 4 in *S. aureus* DnaK complexed with O (OSaD) (Fig. 3C). Moreover, both the E complexes of SaD (ESaD) and PaD (EPaD) have maintained the total number of hydrogen bonds at around 1. However, in CSaD, a sharp decline in the number of intermolecular hydrogen bonds can be observed at 5 ns time point from around 4 to between 0 and 1, which can explain the escape of ligand from its initial docking pocket. Stable active residues have been observed in CPaD (LYS 70, GLU 171, GLU 267), EPaD (ARG 345), and OPaD (THR 11, ASP 194) complexes, as well as in OSaD (GLY 312) complex (Fig. 3G).

In all MD simulation systems, the root-mean-square fluctuations of DnaK receptor maintained at around 0.5 nm (5 Å) except for the C-terminal end where the disordered regions were localized (Fig. 3E). Besides, the RMSD of C and E maintained at 0.5 nm in PaD receptor, while higher RMSD values of around 0.8 nm have been observed in CSaD and ESaD (Fig. 3D). The ligand RMSD were relatively lower in O, compared to others, for which RMSD of 0.4 nm was reported in both SaD and PaD cases. The interaction energies of all systems maintained stably throughout the simulation, except for CSaD complex of which a sharp decrease of Coulomb potential can be observed at around 6 ns time point (Fig. 3F). In general, E maintained the lowest total interaction energies, followed by C and O (Fig. 3H).

**Discussion**

Over the years, the commendable development in the field of virtual screening has enabled time- and cost-efficient drug discovery along with repurposing. Herein, we have carried out a scaffolded approach to antimicrobial drug discovery from yellow variety of Malaysian *N. lappaceum* L. fruit epicarp crude extracts. The first upstream set of experimental work comprised the extraction of the plant product, followed by characterization of their antimicrobial property and chromatographic identification of chemical compounds from therein. This was coupled with downstream set of computational analyses comprising virtual screening and pharmacological predictions of extracted chemical compounds against potential drug targets. To this end, molecular dynamic simulation has taken a step forward to uncover new potent bioactive compounds which can target both gram negative and positive bacteria at the same time. Our study delineates a method to uncover potent chemicals which might have contributed in the antibacterial activities of plant products like *Nephelium lappaceum* epicarp, to be further utilized for drug discovery, repurposing, or other *ab initio* synthetic enhancements.

Extraction is the key stage to obtain the diverse bioactive chemical compounds from plant products. These chemical determinants display different solubility with different organic solvents such that a screening with different solvents helps to bring forth the best one for further exploration. Thus, in order
to explore the extraction of biologically active constituents, several organic solvents were utilized in our study. This was initiated with a sequential extraction process of utilizing solvents like chloroform (CF), ethyl acetate (EA), acetone (AC), ethanol (ET), methanol (MT) and water (WT), in order of their increasing polarity. Our study revealed that the yellow variety of Malaysian *N. lappaceum* epicarp crude extracts exhibited varied inhibitory activities against the six tested MDR pathogens, namely, *B. subtilis*, methicillin-resistant *S. aureus* (MRSA), *S. pyogenes*, *P. aeruginosa*, *K. pneumoniae* and *S. enterica*. In fact, the EA and AC fractions from the sequential mode remarkably inhibited about all the Gram-positive and Gram-negative tested pathogens (TP) while the remaining solvent fractions responded moderately or poorly, thereby providing a strong clue to proceed for further direct extraction from these solvents. Similar results were obtained from the direct extract fractions of EA and AC showing the least MIC value of 250 µg/mL against the TP. For a double cross-checking, the remaining direct fractions did not inhibit the TP even at the highest concentration of 2000 µg/mL. Despite similar reports to our findings by Mohamed *et al.*[^33^]; Thitilertdecha *et al.*[^34^] and Tadtong *et al.*[^35^], a comprehensive chemical profiling to unearth plausible determinants potential enough against the MDR pathogens is lacking till date.

Based on the prominent antibacterial effects of the EA and AC extracts of *N. lappaceum* fruit epicarp, we perceived that both fractions could harbor important bioactive molecules. Thus, we subjected the sequential extracts of EA and AC fractions to HPLC analysis. The results confirmed the presence of some important phenolic compounds with antioxidant properties namely, malic acid, vitamin C, chlorogenic acid, epigallocatechin gallate, quercetin and catechin hydrate (Table 3). Notably, ethyl acetate and acetone have been found to be the competent solvent to extract total flavonoid and phenolic compounds[^32^]. Nazir *et al.*[^36^],[^37^], however, reported the aforementioned compounds in various other organic solvent extracts of *Silybum marianum* and *Elaeagnus umbellate*.

To this end, an extensive spectrum of chemical classes was revealed after LC-MS analysis and included, terpenes, alkaloids, polyunsaturated and monounsaturated fatty acids among others, that were present in both the extracts. Among these, only 21 of the 54 compounds (with above 86% MFG scores) of the EA fractions have been reported to possess antibacterial activities (Table S3). Interestingly, most of them have been newly reported (within the last five years) including L2[^38^], L3[^39^], L6[^40^], L10[^41^], L12[^42^], L16[^43^], L20[^44^], L22[^45^], L26[^46^], L28[^47^], L31[^48^], L46[^49^], L48[^50^], L51[^51^], L53[^52^] and L54[^53^]. Others are known for some time, namely, L7[^41^], L21[^46^], L35[^51^], L37[^52^] and L45[^53^]. Similarly, the AC extract fractions contained only 10 from 44 compounds with reported antibacterial activities (Table S4). Of these, except for L24[^51^] and L35[^53^], all were reported recently. These included L2[^40^], L5[^58^], L10[^44^], L13[^48^], L19[^50^], L36[^54^], L38[^55^] and L42[^57^]. Thus, possibilities exist for those 35 and 26 compounds from EA and AC fractions, respectively, with no matched identity with the library, (Table S5, S6), to be medically important, though, further characterization is required to evaluate their usages.

Further revelation of important biomolecules was authenticated through GC-MS (Table 4) besides the HPLC and LC-MS analyses mentioned above in Tables 3 & S3-S4, respectively. Notably, both sequential and direct extracts of EA and AC fractions were analysed through GC-MS. Unlike the LC-MS reported...
compounds, however, about 50% of the chemical components, unearthed through GC-MS, are unknown for their antibacterial activity. For instance, in case of sequential extracts, for both the EA and AC fractions, only 4 molecules were detected (with area % scores above 0.5%) without any a priori antibacterial activity (Table 4). Likewise, for the direct extract fractions of EA, only 5 out of 8 compounds detected (with area % scores above 0.5%) were known to possess such activity. These are DGEA1\textsuperscript{19}, DGEA2\textsuperscript{20}, DGEA3\textsuperscript{21}, DGEA4\textsuperscript{22} and DGEA5\textsuperscript{23}. However, for the AC extract fraction, 4 out of total 6 compounds detected, were reported to possess the antibacterial effect. These are DGAC2\textsuperscript{19}, DGAC3\textsuperscript{24}, DGAC4\textsuperscript{20} and DGAC6\textsuperscript{21}.

With a set of 91 compounds obtained through chromatographic analyses, we have conducted a computational analysis for virtual screening through molecular docking to short list a selective set of chemicals via pharmacokinetics consideration (Table S9). Pharmacokinetics is an important criterion when it comes to drug discovery and drug design, especially pertaining to bioavailability and toxicity. Herein, we have considered several parameters for absorption, metabolism, drug-likeness, and bioavailability for selecting the ideal drug for potential pharmacological application in the future. For instance, good GI absorption can allow absorption into bloodstream during oral consumption, while bad blood-brain barrier (BBB) permeability can avoid interruption to the central nervous system\textsuperscript{59}. P-glycoprotein (PGP) substrates are being actively effluxed from the cells thereby resulting in low absorption into the blood circulation\textsuperscript{60,61}. Besides these, cytochrome P450 enzymes are crucial in the metabolism of most clinical drugs. Hence, the inhibition of cytochrome P450 enzymes can lead to decreased drug metabolism and possibly, adverse health complications, due to drug-drug interaction upon co-prescription with other drugs\textsuperscript{62,63}. Moreover, the drug-likeness rules, such as the Lipinski rule of five, work by predicting pharmacological behavior upon oral administration based on the chemical properties of potential drugs\textsuperscript{64}. Lastly, bioavailability takes consideration of both absorption and distribution of the drugs, of which the eventual presence in blood circulation upon oral consumption is evaluated.

DnaK protein belongs to the 70 kDa heat shock protein (HSP70) family, which functions as molecular chaperone mediated by its ATPase activities\textsuperscript{65}. DnaK protein has been reported to be central in mediating bacterial stress responses. Among these, increase in antimicrobial susceptibilities and decrease in survivability in host, have been manifested in DnaK mutants\textsuperscript{66–68}. Moreover, our previous work on whole-genome analysis (WGA) of protein interaction network (PIN) reported that DnaK protein was crucial in mediating quorum sensing in multidrug resistant \textit{Proteus mirabilis}\textsuperscript{69}. Furthermore, WGA analyses of PIN from MDR pathogens like \textit{P. aeruginosa}, \textit{S. aureus}, \textit{S. enterica}, \textit{S. pneumoniae}, \textit{P. mirabilis}, \textit{Acinetobacter baumannii}, \textit{Escherichia coli} and \textit{Mycobacterium tuberculosis} revealed DnaK to be among the top 10 crucial proteins indispensable for the cellular integrity of the bacteria\textsuperscript{70}. Also, the ATP-binding pocket of DnaK chaperone has been indicated to be druggable and shown promise to cope with MDR in both gram negatives and positives (Tan & Lahiri, unpublished data). Hence, DnaK protein has been selected for the \textit{in-silico} study, herein, as a promising drug target for MDR bacteria by inhibiting its ATP binding pocket, which can result in its impairment of chaperone function.
Through our computational screening of the chemical libraries of the *N. lappaceum* L. fruit epicarp extractions, Catechin (C), Eplerenone (E), and Oritin-4-beta-ol (O) were shortlisted as the promising antimicrobials in combating the MDR pathogens by dint of their capacity in targeting the DnaK protein and having good pharmacological profiles. Despite being PGP substrate, C has manifested strong binding affinity to DnaK and therefore, can result in effective DnaK functional inhibition with a small amount. Otherwise, PGP inhibitors like C can be co-prescribed easily as it has a good metabolic profile. Moreover, C has been well-characterized for its antibacterial activities and known for its ability to cause leakage of bacterial cellular contents along with increased intracellular reactive oxygen species production in both gram negatives and positives\(^\text{71,72}\). However, the biological targets of C have not been described. As DnaK protein is crucial in bacterial stress response, by inhibiting the DnaK chaperone function, the bacterial cellular and biomolecular integrity can be affected upon receiving environmental oxidative stress. Herein, we showed that in *P. aeruginosa*, C could bind stably to the ATP-binding pocket of DnaK throughout the MD simulation with 3 stable active residues (LYS 70, GLU 171, and GLU 267), while maintaining the ATP-bound conformation of the DnaK protein without the necessity for ATP binding (Fig. 2B, 3A). This reflected the inability of the ATP molecules to bind the CPaD (Catechin-bound DnaK protein of *P. aeruginosa*) as also a complete halting of the normal DnaK chaperone function via conformational changes ensuing ATP hydrolysis. However, C could not inhibit SaD (DnaK of *S. aureus*) the same way, due to its inability to maintain the integrity of NBD domain and thereby escaping from the binding pocket. It is this binding pocket which allows subsequent binding of ATP molecules on DnaK to continue the chaperone function. On the contrary, herein, we present the discovery of two novel potential compounds, E and O, whose antibacterial activities have not been reported and/or described earlier. Notably, E has been widely utilized in cardiovascular implications and as diuretics\(^\text{73,48}\). O, however, has not been explored to confer any biological significance. Despite that, it is notable that the chemical structure of O is analogous to C (Fig. 1), with the sites of hydroxylation being slightly different.

Throughout the molecular dynamics simulation (MDS) processes, only 1 or 2 hydrogen bonds can be observed in EPaD and ESaD, which suggested weak protein-ligand electrostatic interactions. This can be explained by the chemical structure of E, being crowded with carbonyls and ethers which are weak bases, and hydroxyl groups are lacking. The ligand, however, has been retained in the docking pocket over the course of MDS. This probably suggests that hydrophobic (van der Waals) interactions were dominant in this case. This was reflected through the intermolecular interaction energies (Fig. 3F), of which the Lennard-Jones potentials were much higher than Coulomb potentials in Eplerenone-DnaK (ED) complexes, while the reverse was observed in for C and O. Moreover, the binding conformation of E in PaD did not “cover up” completely at the binding site of phosphate groups of the ATP for which further wet-lab confirmation is required. Furthermore, among the three ligands simulated, O manifested the best binding capabilities to both PaD and SaD with rich intermolecular electrostatic interactions and highest total interaction energies. After MDS, the active residues THR 11 and ASP 194 were retained in OPaD, while GLY 312 was retained in OSaD. Again, despite being structurally analogous to C, O manifested good predicted pharmacological properties in all the aspects considered. Therefore, with better binding capabilities to DnaK receptor and pharmacological properties, herein we report O to be a more potent
antibacterial compound compared to the well-known C, which is active against both the gram positive and negative bacteria.

Conclusion

Our findings reindicate the promising antibacterial effects, of the yellow variety of Malaysian Rambutan (*N. lappaceum L.*) fruit epicarp crude extracts, against selected Gram-positive and Gram-negative MDR pathogens. In this context, particularly ethyl acetate and acetone (sequential and direct) extracts demonstrated remarkable antibacterial effects toward all tasted pathogens, while remaining fractions including, chloroform, ethanol, methanol and water did not exhibit such potential. Nevertheless, *N. lappaceum* presents itself to be a novel source for antibacterial compounds with high potential for the development of pharmaceutically valuable drugs. Further studies are mandatory to separate the specific compound(s) responsible for the desired effects, and to develop our knowledge on the other unseen potentials in *N. lappaceum*.

Materials And Methods

Solvents.

All solvents, used for preparation of crude extractions, were of HPLC grades In order of increasing polarities, these were Chloroform (99.9%, Sigma-Aldrich, LiChrosolv, Malaysia), Ethyl acetate, Acetone (99.5% Chemiz, Malaysia), Ethanol, Methanol (99.8%, ChemAR, Systerm, Malaysia) and double distilled Milli-Q Type 1 water (MilliporeMerck, Germany). Solvents used for LC-MS and GC-MS were of MS grades.

Plant Product.

The yellow variety fruits of *N. lappaceum L.* were purchased from local marketplace, Bandar Sunway, Selangor, Malaysia.

Tested microorganisms.

Six clinical isolates, used in the study, were obtained from the Department of Biological Sciences, Sunway University, Malaysia. These were *Streptococcus pyogenes* (ATCC-49399), *Bacillus subtilis* (ATCC-11774), methicillin-resistant *Staphylococcus aureus* (MRSA) (MTCC-381123), *Pseudomonas aeruginosa* (ATCC-10145), *Klebsiella pneumoniae* (ATCC-700603) and *Salmonella enterica* (ATCC-14028). All strains were tested to be multidrug resistant.

Preparation of crude extracts.
Epicarp crude extracts were prepared following the method of Do et al.\textsuperscript{75} using the solvents mentioned earlier for the direct extracts. For the sequential method of extraction, the mentioned solvents were used in order of increasing polarity \textit{viz} chloroform<ethyl acetate<acetone<ethanol<methanol<water. In both the cases, essentially, the peels of \textit{N. lappaceum} were removed from the fruit and washed thoroughly with running, followed by, distilled water to remove contaminants and thereafter dried using freeze-dryer. Dried peels were ground into fine powder using an electric grinder. To produce different fractions of crude extracts, 10 g of powder was extracted in 100 mL of selected solvents. The solution was mixed thoroughly by using incubator shaker (Yihder LM-530D Incubator Shaker, Taiwan) for 24 h. To separate supernatant, the solution was centrifuged (Eppendorf 5810 R Centrifuge, Germany) at 4000 rpm for 10 min at 4 °C to completely eliminate the leftover fine sediments. The solvent extracts were concentrated using Rotary evaporator, and further with vacuum concentrator until a viscous extract was obtained. All extracts were stored at 4°C for future experiments.

**Potential \textit{in-vitro} antibacterial activities of yellow rambutan fruit epicarp extracts.**

**Disc Diffusion assay.**

Seed culture of the tested pathogen was consistently swabbed on agar plate. Sterilized blank paper discs were separately impregnated by different concentration of extracts (250 to 2000µg/ml) and placed on agar plate. The plates were incubated at 37°C for 16h. The antibacterial activity was noted by measuring the diameter of inhibition zone. Gentamicin (10µg/disc) was used as positive control while DMSO (<1%) was kept as negative control. All the experiments had technical triplicates and were performed twice to render two biological replicates.

**Broth Dilution Assay.**

A broth micro-dilution method was used to evaluate the minimum inhibitory concentration (MIC) values of crude extracts using Clinical & Laboratory Standards Institute (CLSI) procedures. Each extract (5μL) was added into the wells of a 96 well plate comprising $10^5$ CFU/mL bacterial cells. The 96 well plates were incubated at 37°C for 16 h. Final concentrations ranged from 250 to 2000 µg/mL. Three controls comprising, gentamicin 10µg/mL (positive control), DMSO <1% (solvent control) and bacterial inoculum (negative control) were included in each test. The lowest concentration of the tested extract showing inhibitory effect against the pathogens, recorded via the Microplate reader (TECAN, Infinite-M200-PRO), was taken as the MIC value. All tests, having technical triplicates, were confirmed twice. Both the fractions of ethyl acetate and acetone extracts gave promising results with which all chromatographic analyses were carried out.
Exploration of chemical constituents through chromatographic analyses.

**High performance Liquid Chromatography (HPLC).**

Ethyl acetate and acetone extracts were used as samples for qualitative phytochemical screening via HPLC using Agilent-1260 infinity system, according to the reported method of Zeb\(^76\). Briefly, one-gram sample extract was mixed in methanol and water (1:1; 20 mL; v/v) and heated at 70°C for 1 hour in water bath. This was centrifuged at 4000 rpm for 10 minutes and 2 mL of the supematant was filtered into HPLC vials through Whatman filter paper. The separation was performed via Agilent-Zorbax-Eclipse column (XDB-C18). Column gradients system comprised solvent B and C. Solvent B consisted of deionized water: methanol: acetic acid having a ratio of 180: 100: 20; v/v while solvent C had deionized water: methanol: acetic acid in the ratio of 80: 900: 20; v/v. Gradient system was started by solvent B for 100%, 85%, 50% and 30% at 0, 5, 20 and 25 minutes followed by solvent C (100%) from 30-40 minutes. Elution occurred after 25 minutes. The ultraviolet array detector (UVAD) was set at 280 nm for the antioxidants analysis and chromatogram were documented using retention times. UV spectra of compounds and accessible standards along with quantification was carried out by taking the percent peak area. Quantification of the antioxidants was measured by formula:

\[
C_x = \frac{A_x \times C_s (\mu g/ml) \times V (ml)}{A_s \times Sample (wt.in g)}
\]

\(C_x\)= Sample concentration; \(A_s\)= Standard peak area; \(A_x\)= Sample peak area; \(C_s\)= Standard concentration (0.09 µg/ml).

**Liquid Chromatography and Mass Spectrometry (LC-MS).**

A mixture of standards and new metabolites found in the ethyl acetate and acetone fractions were analyzed via LC-MS exactly as per the method reported by Yap et al.\(^77\). In order to eradicate systematic errors, reference solution was used with the two ions, having m/z of 121.0508 and 92266.0097, being selected for mass calibration. Finally, the mass spectra for the compounds present in ethyl acetate (EA) and acetone (AC) fractions were run against the database of NIST (National Institutes of Standard and Technology, Gaithersburg, MD, USA) Mass Spectral Search Program-2009 version 2 for the documentation of homologous compounds over Agilent Mass-Hunter Qualitative Analysis B.05.00 software.
**Gas chromatography–mass spectrometry (GC-MS).**

Ethyl acetate and Acetone fractions were subjected to gas chromatography-mass spectrometry (GC-MS) analysis, using Agilent technologies model 7890B GC System coupled with Pegasus HT High Throughput TOFMS (Leco Corp., MI, USA). An aliquot of an extract of 1ml was injected to the GC-MS apparatus. Next, Agilent J&W HP-5MS (phenyl methyl siloxane, length 30 m, Dia. 0.32 mm, Film, 0.25µm) analytic column was used to separate components under an inert atmosphere of helium (1.5 mL/min). Other standardized parameters utilized during the process: oven temperature of 80°C (2 min) was increased to a temperature of 300°C at the rate of 3°C/min, solvent delay time was 5 min, inlet line temperature was 225°C, and ion source temperature was 250°C. Mass spectra were taken at 70 eV and acquisition mode-scan was 20-1000 amu while sixty-four (64) minutes was the GC run time. The interpretation of mass spectrum and documentation of phytochemicals present in the fractions were achieved via the database of NIST libraries.

**Virtual Screening of Chemical determinants from chromatographic analyses.**

**In silico Protein Model Generation.**

*S. aureus* (Sa) and *P. aeruginosa* (Pa) were chosen as gram-positive and gram-negative bacterial representatives for computational analyses of DnaK protein binding. 3D structures of DnaK proteins, from the aforesaid species, were generated via homology modelling using MODELLER version 9.24. DnaK has two conformations, namely, the open or ATP-bound and the closed or ADP-bound conformation. Herein, we focused on the open conformation of DnaK, to identify potential competitive inhibitors of ATP in order to prevent proper functioning of DnaK protein.

The protein sequences of *Sa* and *Pa* DnaK were obtained from UniProtKB with accession IDs of Q2FXZ2 and A6VCL8 (UniProt Consortium, 2019), respectively. To search for suitable homology modelling templates, both NCBI BLASTp and the MODELLER in-built build_profile.py were utilized. For *Pa* DnaK (PaD), the templates were full-length ATP-bound *E. coli* (Ec) DnaK protein structures (PDB ID: 5NRO, Chain: A, Query Coverage (QC): 94%, Percent Identity (PI): 79.50%, Resolution (R): 3.25 Å; PDB ID: 4JNE, Chain: A, QC: 94%, PI: 78.80%, R: 1.96 Å; and PDB ID: 4B9Q, Chain: A, QC: 94%, PI: 77.96%, R: 2.40 Å). For *Sa* DnaK (SaD), besides the aforementioned *Ec* DnaK (EcD) models, one additional template, from *Geobacillus kaustophilus* DnaK protein (PDB ID: 2V7Y, Chain: A), was selected due to the high percentage of sequence identity expected as per the gram positive character of *S. aureus* and *G. kaustophilus*. As this template structure was in closed conformation and we were only interested in the open conformation, only the Nucleotide Binding Domain (NBD, residues 1 to 350 in template model) which does not differ much in both conformations, were taken into consideration for homology modelling, and the remaining C-terminal residues modelling were guided by the *Ec* models to shape an open conformation. Therefore, the
templates for SaD were (PDB ID: 2V7Y, Chain: A, Template Residues: 1-350, QC: 57%, PI: 83.19%, R: 2.37 Å; PDB ID: 5NRO, Chain: A, QC: 93%, PI: 56.19%, R: 3.25 Å; PDB ID: 4JNE, Chain: A, QC: 92%, PI: 55.54%, R: 1.96 Å; and PDB ID: 4B9Q, Chain: A, QC: 94%, PI: 55.43%, R: 2.40 Å). 5 homology models were generated for each protein of SaD and PaD, and the models with lowest DOPE (discrete optimized protein energy) scores were selected for downstream virtual screening for both. The SaD and PaD homology models were validated via Swiss-Model Structure Assessment and SAVES v5.0 servers\(^8\) (Table S7).

**Druggable Pocket Validation.**

To validate the druggability of the ATP docking pocket, we have conducted ligand binding site prediction using P2Rank from PrankWeb server\(^8\). P2Rank predicts the chemical druggability on protein solvent-accessible surface via a non-templated machine learning approach. The ATP binding pocket was predicted to be druggable and ranked first in both cases of SaD and PaD (Table S8; Fig. S8). These pockets from SaD and PaD were further considered to be targeted for virtual screening.

**Molecular Docking with Chemical Determinants.**

POAP pipeline, Samdani & Vetrivel\(^8\) was followed for an *in silico* virtual screening of the chemical compounds obtained through different chromatographic separation. SMILES notations of these compounds were obtained and their 3D models (in mol2 format) were generated through POAP Ligand Preparation pipeline. To this end, Chimera was utilized to generate physiological protonation states of ligands, and PDBQT files were prepared\(^8\). Ligand optimizations were carried out via POAP Ligand Preparation pipeline utilizing MMFF94 force field, being optimized for drug-like organic molecules and molecular docking\(^8\). Out of the 50 conformers, generated for each ligand through Weighted Rotor Search approach, only the best conformers were retained. Finally, the ligands were subjected to energy minimization for 5000 steps by the conjugate algorithm.

The macromolecule receptors, pertaining to the SaD and PaD proteins, were prepared using AutoDockTools. AutoDock 4.2, aided by POAP pipeline, was utilized for the virtual screening process\(^8\). For AutoDock parameters, 100 generations of Lamarckian Genetic Algorithm were set for each protein-ligand complex. To fit in the previously predicted pocket, docking grids were adjusted into squares of 24 Å with x, y, z coordinates of 17.647, 75.43, 27.766, and 18.069, 74.299, 28.532, for SaD and PaD, respectively. For the silicon-containing compound among the set of ligands, molecular docking was separately carried out with AD4.1_bound parameter file, obtained from AutoDock, wherein parameters for silicon atoms (Rii=4.3; eii=0.402) were added\(^8\).

**Pharmacological Properties Screening.**
Pharmacological properties, encompassing pharmacokinetics, drug likeness, and molecular information for each chemical compound, were predicted using SwissADME\textsuperscript{87}.

**Molecular Dynamics Simulation.**

Ensuing virtual and pharmacological screenings, potential drug candidates were rationally selected to undergo molecular dynamics (MD) simulation via GROMACS version 2019.3\textsuperscript{88}. CHARMM36 force field (Version July, 2020), along with TIP3P water model, was utilized for macromolecule processing\textsuperscript{89}. Avogadro software was utilized for mol2 format conversion and complete protonation (protonation of non-polar atoms)\textsuperscript{90}. The Perl script, sort_mol2_bonds.pl, written by Justin Lemkul was utilized for bond order arrangements in ligand mol2 files. Then, topologies of the ligand models were generated through CGenFF server, and a python script (cgenff_charmm2gmx.py) was utilized to convert topologies for CHARMM to GROMACS\textsuperscript{91}. Solvation was carried out in a dodecahedron box ranged 1.0 Å from the protein-ligand complex. The system was then being ionized to achieve electrostatic neutralization. Subsequently, the system was subjected to energy minimization via steepest descent algorithm until convergence at maximum force of less than 1000 kJ mol\textsuperscript{-1} nm\textsuperscript{-1} (Fig. S9). Potential energy shifts of the systems were monitored herein.

Equilibration of the systems were carried out via NVT and NPT ensembles for 50000 steps (100ps), with temperature, pressure, and density shifts being monitored therein. Subsequently, production MD simulations were carried out for 5000000 steps (10ns) to observe protein-ligand interactions. RMSD (Root Mean Square Deviation) values of ligands and receptors, number of hydrogen bonds between ligands and receptors, and ligand-receptor interaction energies (Coulombic interaction energies and Lennard-Jones energies) were computed throughout the MD simulations. Total interaction energies were computed, and errors were estimated via error propagation by addition.

**Declarations**

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**Author contributions**

CL conceived the concepts, planned and designed the analyses. AA extracted the natural products and assessed their antibacterial activities. MZ conducted and analyzed HPLC, helped by AA. SA conducted LC-MS with the results being analyzed by AA and CL. YYY provided the plant products and co-supervisory
inputs along with EK. YCT conducted the computational part with occasional inputs from CL. Concepts for artwork were generated and executed by AA and TCY with guidance provided by CL. AA and YCT primarily wrote the manuscript aided by editorial upgradation by CL.

**Additional Information**

The authors declare that they have no conflict of interest, financial or otherwise, for this research.

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**Figures**

![Catechin](image1.png)

**Catechin**

![Oritin-4-beta-ol](image2.png)

**Oritin-4-beta-ol**

![Eplerenone](image3.png)

**Eplerenone**
Figure 1

Chemical Structure of Catechin, Oritin-4-beta-ol, and Eplerenone.