INTRODUCTION

Several already drugs were discovered from the natural products. Especially, the treatments of infectious diseases and oncology have benefited from numerous drugs which were found in natural product sources. Some of new and interesting natural compounds with biological activities have been published in the past few years. The potent biological activities, no side effects, and economic viability of medicinal properties from natural products have been investigated in the recent scientific developments worldwide. Bioactive compounds from medicinal plants showed pharmacological or toxicological effects in man and animals. The typical bioactive compounds are secondary metabolites including steroids, glycosides, phenolics, tannins, anthocyanins, flavonoids, and alkaloids. The basic structural of alkaloids are nitrogen-containing compounds. The several properties of alkaloids are bitter taste and usually with potent biological activities. Moreover, these secondary metabolites have diverse clinical biological properties [1-3]. Name of the genus Tabernaemontana is the birthplace (Tabernaemontanus) of J. Th. Müller, a German physician and botanist who was born in Bergzabern and died in Heidelberg in the Pfalz in 1590. The genus of Tabernaemontana belonging to the Apocynaceae family and composing about 100 species distributed throughout the tropical and some subtropical parts of the world. Many of plants in the Tabernaemontana species are used in traditional medicine and for other purposes for the treatment of sore throat, hypertension, and abdominal pain [4-6]. Plants of Tabernaemontana genus are a prolific source of the monoterpene indole alkaloids and they have been shown to produce many skeletal types such as secatabersonine alkaloids, bis-vobtusine-type alkaloids, and bis-vobasinyl-ibogan indole alkaloids [7-9]. These alkaloids are a diverse class of natural products, comprising over 2000 members and possess a range of chemical structures and a wealth of biological activities including anticancer, antimalarial, and anti-arrhythmic agents [10]. Therefore, the biogenesis, classification, and biological activities of the indole alkaloids found in Tabernaemontana species were discussed in this review and its bring the research up-to-date on the bioactive compounds produced by Tabernaemontana species, directly or indirectly related to human health.

Classification and biogenesis of monoterpene indole alkaloids

Many skeletal types of monoterpene indole alkaloids from Tabernaemontana plants were exhibited a wide array of biological activities. Then, the biosynthetic pathways of some classes of these alkaloids are investigated. All monoterpene indole alkaloids are derived from aromatic amino acid tryptophan and the iridoid terpene secologandin (Scheme 1). Tryptophan converts to tryptamine using tryptophan decarboxylase which is a pyridoxal-dependent enzyme. The specific iridoid precursor was subsequently identified as secologandin. After that, tryptamine was reacted with the secologandin using the enzyme strictosidine synthase catalyzes a stereoselective Pictet-Spengler condensation to yield strictosidine (iso-vincoside) (5 stereochemistry at C5, Scheme 1), and this step is utilized in the first committed step of monoterpene indole alkaloid biosynthesis. The Rubiaceae, Nyssaceae, Loganiaceae, and Apocynaceae families of medicinal plants each produce monoterpene indole alkaloids with dramatically diverse structures. The mechanisms and control of the processes by which strictosidine rearranges into these diverse families of products remain one of the most fascinating problems in secondary metabolism [10].

The biosynthesis pathway of strictosidine rearranges also use as a basis for the classification of the monoterpene indole alkaloids occurring in the genus Tabernaemontana. Eleven classes of these alkaloids were reported to the structural characteristics of their skeletons including (1) vincosan, (2) corynanthean, (3) vallesiachotaman, (4) strychnan, (5) aspidospermatan, (6) plumeran, (7) eburnan, (8) ibogan, (9) tacaman, (10) bis-indole alkaloids, and (11) miscellaneous. The classification of the monoterpene indole alkaloids occurring in the genus Tabernaemontana were shown in Table 1 and Scheme 2 [4,10].

Bioactive compounds from the genus Tabernaemontana

The plants of genus Tabernaemontana have been used in traditional medicine for the treatment of sore throat, hypertension, and abdominal pain [6]. The chemical constituents from several parts of the Tabernaemontana species were reported as monoterpene indole alkaloid compounds, and these compounds were shown to exhibit a wide array of biological activities including anticancer, antimalarial, and anti-arrhythmic agents [10]. This review brings the research up-to-date on the bioactive compounds produced by Tabernaemontana species, directly or indirectly related to human health.
Ibogaine (6), ibogaline (7), desethyl-voacangine (8), vochalotine (9), and affinsine (10) from the chloroform extract of stalk of *T. australis* (Muell. Arg) Miers by gas chromatography-mass spectrometry (GC-MS) (Scheme 3). Some of them showed qualitative anticholinesterase activity using thin-layer chromatography assay and physostigmine and galanthamine were used as positive drugs [11].

**Tabernaemontana catharinensis** A. DC.

In 2007, Soares et al. reported the extraction of branches and leaves of *T. catharinensis* by supercritical fluid using a mixture of CO₂ plus ethanol (supercritical fluid extraction). It was found that the enrich indole alkaloid fraction was showed a strong active against *Leishmania amazonensis* (leishmaniasis). Moreover, this fraction was inhibited NO production induced by IFN-γ plus lipopolysaccharide which some of effect for antiparasite activity and they showed no cytotoxicity with host cells [12].

In 2013, Boligon et al. reported that the leaves extract of *T. catharinensis* were evaluated total phenolic, flavonoid, tannin, and alkaloid contents. In addition, this extract was tested the antioxidant activities using 2,2-diphenyl-1-picrylhydrazyl free radical and thiobarbituric acid-reaction species methods. From these results, the ethyl acetate (EA) fraction of leaves extract was the most effective fraction against two reactive species and showed high total phenols, flavonoids, tannins, and alkaloids contents. The results indicate that the leaves extracts of *T. catharinensis* has antioxidant potential and can be a promising source of natural antioxidants [2].

In 2015, the same group reported that the crude extracts, fractions, and subfractions of *T. catharinensis* were evaluated antimycobacterial, antimicrobial, and antiviral activities. The dichloromethane fractions showed strong active against *Enterococcus faecalis*, *Micrococcus sp.*, *Staphylococcus aureus*, and *Bacillus subtilis* with minimum inhibitory concentration (MIC) values of 3.125–1000 µg/mL. Considering the Gram-negative bacteria, only butanolic fraction was effective against *Proteus mirabilis* and *Aeronomas sp.* (MIC=62.5 µg/mL and 250 µg/mL, respectively). In addition, the fungi *Candida albicans*, *Candida glabrata*, *Cryptococcus neoformans*, *Saccharomyces cerevisiae*, *Aspergillus flavus*, and *Aspergillus fumigatus* were particularly vulnerable for dichloromethane fraction (MIC=31.25–1000 µg/mL). The fractions and subfractions were effective against *Mycobacterium smegmatis* (MIC=19.53–156.25 µg/mL). Dichloromethane (selectivity index=0.3=77.92), EA (SI=40.27) and NB (SI=28.97) fractions from the leaves exhibited a potential antiviral activity toward Herpes Simplex Virus Type 1 whereas dichloromethane subfraction from the leaves (SI=12.28) and alkaloidal fraction (10.71) maintained this good activity. From these result studies were obtained *T. catharinensis* as a source for new antiviral and antimicrobial therapy [13].

**Tabernaemontana citrifolia** Linn.

In 2010, Marie-Magdeleine et al. reported that three extracts (aqueous, methanolic, and dichloromethane) of *T. citrifolia* fruits, leaves, and root were evaluated in vitro effect against *Haemonchus contortus* on four developmental stages of the parasite including egg hatch, larval development, L3 migration, and adult worm motility. From the results, the different parts of *T. citrifolia* showed significant effects with differences depending on the parasitic stage such as efficacies on the larval development stage for fruits extract. These results suggest that *T. citrifolia* will be to possess anthelmintic activity against *H. contortus* [14].

**Tabernaemontana coronaria** (L.) R. Br.

In 2012, Poomnina et al. evaluated that the acute and subacute toxicological of ethanol extract from *T. coronaria*. The ethanol extract showed no significant effect on hematological lipid and renal profiles and showed only a slight increase in alkaline phosphatase (ALP) and

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**Table 1: Classification of the terpene indole alkaloids from the genus Tabernaemontana**

<table>
<thead>
<tr>
<th>Class</th>
<th>Abbreviation</th>
<th>Structure characteristics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vinosan</td>
<td>D</td>
<td>C (2)-C (3)-C (14) unit, no N (4)-C (17) or N (4)-C (21) bond</td>
</tr>
<tr>
<td>Corynanthean</td>
<td>C</td>
<td>C (2)-C (3)-C (14) unit, N (4)-C (21) bond</td>
</tr>
<tr>
<td>Vallesiachotaman</td>
<td>V</td>
<td>C (2)-C (3)-C (14) unit, N (4)-C (17) bond</td>
</tr>
<tr>
<td>Strychnan</td>
<td>S</td>
<td>C (2)-C (16)-C (15) unit, C (3)-C (7) bond</td>
</tr>
<tr>
<td>Aspidospermatan</td>
<td>A</td>
<td>C (2)-C (16)-C (15) unit, no C (3)-C (7) bond</td>
</tr>
<tr>
<td>Plumeran</td>
<td>P</td>
<td>C (2)-C (16)-C (17)-C (20) unit</td>
</tr>
<tr>
<td>Eburan</td>
<td>E</td>
<td>N (1)-C (16)-C (17)-C (20) unit</td>
</tr>
<tr>
<td>Tacaman</td>
<td>T</td>
<td>N (1)-C (16)-C (17)-C (14) unit</td>
</tr>
<tr>
<td>Ibogan</td>
<td>I</td>
<td>C (2)-C (16)-C (17)-C (14) unit</td>
</tr>
<tr>
<td>Bis-indole</td>
<td>B</td>
<td>Two indole alkaloids attached to each other</td>
</tr>
</tbody>
</table>
lactate dehydrogenase in rats. In addition, the histopathological study also showed inactive on any abnormalities in the liver and kidney. From these results, *T. coronaria* extract can be used in indigenous system of medicine because this extract showed no acutely toxic to the rats. However, further long-term toxicological studies (chronic study) are needed to establish it as medicine for human consumption [15].

*Tabernaemontana corymbosa* Roxb. ex Wall.

This plant provided several novel molecular skeletons of indole alkaloids (Schemes 3-5) which have been shown to exhibit a wide array of biological activities. Lim and co-worker reported the isolation of novel monoterpane indole alkaloids from the stem bark extract of Malaysian *T. corymbosa* which were collected from a different part in Malaysian, including conolutinine (11) [16] and a mixture of (1S, 2'S)- and (1R, 2'R)-diastereomers of alkaloid-lignan conjugates, conoliferine (12) and isoconoliferine (13) [17]. The structure of 11 showed an unprecedented ring system incorporating a diazaspiro center and fused oxadiazepine–tetrahydrofuran rings [16]. In addition, four novel natural products indole alkaloid–hydroxycinnamyl alcohol conjugates, conomicidines A and B (14 and 15) and diastereomeric isonomicidines A and B (16 and 17) [18], four tetracyclic oxindole alkaloids, 7(R)- and 7(S)-geissoschizol oxindole (18 and 19), 7(R), 16(R)– and 7(S), 16(R)–19(E)-isositsirikine oxindole (20 and 21), together with a new taberpsychine derivative, N(4)-demethyltaberpsychine (22) [19], and two new pentacyclic indole alkaloids was possessed an new rearranged dibenz ring system, lirofolines A and B (23 and 24) [20] were also isolated from this plant. Moreover,

Scheme 3: Bioactive indole alkaloids (1–39) isolated from *Tabernaemontana* plants
these research groups reported that the isolation and purification of seven new cytotoxic Aspidosperma-type alkaloids, jerantinines A-G (25–31) against human KB cells (IC₅₀ < 1 µg/mL) [21]. In addition, the isolation of seven indole alkaloids (32–38) from the stem bark extract of Malaysian *T. corymbosa* which were collected near Taiping, Perak, Malaysia was reported by Sim *et al.* Two of the new vobasine alkaloids (32 and 34) showed appreciable cytotoxicity toward KB cells (IC₅₀ ca. 5.0 µg/mL). The structure of the known *Tabernaemontana* alkaloid tronoharine (39) was revised based on newly acquired nuclear magnetic resonance (NMR) data, as well as X-ray diffraction analysis [22].

Moreover, Lim *et al.* were reported the isolation of two secotabersonine alkaloids, jerantiphyllines A and B (40 and 41) and a tabersonine hydroxyindolenine alkaloid, jerantinine H (42), from the leaves extract of the same plant species, but involving plant material collected from a different location and also reported vincamine alkaloid 43 from the leaves of this plant [8]. In 2014, the isolation of two novel dimeric monoterpen indole alkaloids, bistabercarpamines A (44) and B (45) possessing unprecedented bis-vobasinyl-chippiine-type skeleton from the leaves extract of *T. corymbosa* which were collected from Yunnan Province of China was reported Ma *et al.* In addition, bistabercarpamine A (44) showed moderate cell growth inhibitory activity against HepG-2 cells with IC₅₀ of 38.14 ± 1.1 µM [23] and were also reported the isolation of ten indole alkaloids, tabercarpamines A-J (46-55) from the leaves extract of the same species. In addition, the all isolated compounds were evaluated for cytotoxic activity using 3-(4,5-dimethylthiazol-
2-yl)-2,5-diphenyl tetrazolium bromide assay against MCF-7, HepG2, and SMMC-7721 cell lines. The cytotoxicity results, alkaloid 47 showed significant inhibitory effects against all human cancer cell lines. Moreover, alkaloid 48 inhibited the proliferation of HepG2 cells by inducing apoptosis using the annexin-V/PI double-staining assay [24].

In addition, Zhang et al. reported the isolation and evaluation for cytotoxicity against various human cancer cell lines of three vobasinyl-ibogan type bisindole, tabercorines A–C (56–58), together with 17-acetyl-tabernaecorymbosine A (59) from the twigs and leaves of T/corymbosa. It was found that alkaloids 56 and 59 showed significant inhibitory effects with IC\(_{50}\) values comparable to those of cisplatin [25]. In the recently, Nge et al. reported the isolation and identification of eight monomeric (60-67) and two dimeric (68,69) indole alkaloids from the stem bark of Malayan T. corymbosa. The structure and absolute configurations of all new monoterpene indole alkaloids were completely determined based on spectroscopic data and X-ray diffraction analysis [26].

**Tabernaemontana crassa Benth.**

In 2010, Kuete et al. reported the toxicological activity of the hydroethanol extract from the stem bark of T. crassa (Apocynaceae). This extract showed no a dose-related effect in liver, lungs, and kidneys at 0.5 g/kg body weight when the animals received the extract for 6 weeks daily. The acute toxicity of this plant extract indicated the medium lethal dose (LD\(_{50}\)) of 6.75 g/kg body weight after treatment for 48 h and they showed no effected with body weight, serum ALP, alanine aminotransferase, total bilirubin, direct bilirubin, and creatinine (SCr) at the dose of 6 g/kg. The results indicate that the stem bark extract of T. crassa showed no toxic with animal in several conditions and can be a promising source of new natural product drug [27].

**Tabernaemontana dichotoma Roxb. ex Wall.**

In 2013, Zaima et al. reported the screening of medicinal plants targeting vasorelaxant activity. It was found that the methanol extract from T. dichotoma bark showed vasorelaxant activity on rat aorta. After that, this plant extract was isolated to give eight indole alkaloids (Scheme 5) including 10-methoxyalstonerine (70), 10-methoxyaffinisine (71), lochnerine (72), cathafoline (73), (-)-alstonerine (74), 19,20-dehydro-10-methoxytalcarpine (75), alstonisine (76), and alstonal (77). Among of them, sarpagine type (71 and 72), akuammiline type (73), and macroline oxindole type (76 and 77) showed potent vasorelaxant activity. Furthermore, alkaloids 71 and 76 attribute to the inhibitory effect of voltage-dependent Ca\(^{2+}\) channel and receptor-operated Ca\(^{2+}\) channel. In addition, the major compound...
(71) from *T. dichotoma* bark extract showed hypotensive effect on normotensive rats in vivo [28].

*Tabernaemontana divaricata* (L.) R. Br.

Acetylcholinesterase inhibitors (AChE-Is) are the current pharmacotherapy to use for Alzheimer’s disease (AD). In 2006, Ingkaninan et al. reported the bioassay-guided fractionation of the different parts of *T. divaricata* (flowers, leaves, stems, and roots) on AChE activity using the Ellman’s colorimetric method. It was found that the stem (94.72±2.09%) and root (99.72±0.26%) extracts showed high inhibitory activity while the leaves and flower extracts showed lower activity. From these results, led to the isolation of four AChE inhibitors (Scheme 5) including two new bisindole alkaloids, 19,20-dihydrotabernamine (78) and 19,20-dihydro-ervahanine A (79) together with two known bisindole alkaloids, conodurine (80), and tabernaegantine A (81) from the root extract of this plant [29].

The next year, this group was also reported in vivo the evaluation of AChE inhibitory effects and Fos expression on neuronal activity in the cerebral cortex of the ethanolic extracts from roots of *T. divaricata*. This extract showed strong the neuronal activity with the enhancement of Fos expression and AChE inhibitory effects in the cerebral cortex [30]. Moreover, this group was also reported in 2010 for the effects of *T. divaricate* root extract on β-amyloid25-35 peptides-induced cognitive deficits in mice. After mice were treated with root extract, it was found that this extract improved the memory impairment and attenuated the brain levels of AChE activity induced by Aβ25-35 peptides [31]. On the other hand for AD treatment, Chaiyana et al. reported that the alkaloidal extract from *T. divaricata* stem was showed enhancing the acetylcholine level in Alzheimer’s patients. From these results, they reported the isolation of 3'-R/S-hydroxyvoacamine (82) which showed noncompetitive inhibitor against AChE with an IC₅₀ value of 7.00±1.99 µM [32,33].

For cytotoxic activity of *T. divaricata*, Thind et al. reported to study cytotoxic activities against HCT-15 (Colon), HT-29 (colon), 502713 (colon), MCF-7 (breast), PC-3 (prostrate) cell lines and evaluated the mechanism of cytotoxicity including hydroxyl radical scavenging and topoisomerase inhibitory activities of the leaves extracts (hexane, chloroform, EA and methanol) of this plant. It was observed that the EA extract was effective against only one colon cell line (502713), whereas the chloroform extract was effective against all the three colon cancer cell lines. Moreover, the EA extract also showed selectively inhibition of topo II in topoisomerase II relaxation assay [34]. In 2013, Zhang et al. reported to study of conophylline (83), a bis-indole alkaloid (Scheme 6): Bioactive indole alkaloids (83-95) isolated from *Tabernaemontana* plants.
6) consisting of two pentacyclic aspidosperma skeletons, isolated from *T. divaricata*, which has been found to induce b-cell differentiation in rat pancreatic acinar carcinoma cells and in cultured rat pancreatic tissue [35]. In the same year, Bao *et al.* reported the isolation of five new cytotoxic vobasinyl-ibogan-type bisindole alkaloids (Scheme 6), tabernaricatines A-E (84-88), two new monomers, tabernaricatines F and G (89 and 90), and 24 known indole alkaloids were isolated from the aerial parts of *T. divaricata* [36]. In 2012, two research groups including Jain *et al.* and Mukhram *et al.* reported to evaluate the antifertility effect of leaves and flowers extracts of *T. divaricata* in male rats and female albino mice. In the male rat results, the extracts produced dose-related effect on reproduction. The effect may have an inhibitory influence on gonadotropin release which may be responsible for the decline in testosterone production, leading to change in spermatogenesis. For female albino mice, disturbance on the estradiol secretion with significant decrease during estrous stage of the cycle observed with the extract treatment may be due to impairment in the release of luteinizing hormone and follicle stimulating hormone causing hormonal imbalance. These researches could also suggest the anti-fertility effect of leaves and flowers extracts of *T. divaricata* in male rats and female albino mice. In 2013, Jain *et al.* reported the in vivo anti-inflammatory activity of *T. divaricata* leaves extract in male albino mice. The hexane fraction which is very rich of flavonoid compound showed a very high anti-inflammatory activity. This fraction was also active than positive drug indomethacin [40].

In 2009, Hirasawa *et al.* reported the isolation a novel tetrakis monoterpene indole alkaloid 91, namely, alasmontamine A, from the leaves of this plant and this novel compound exhibited moderate cell growth activity against HL-60 cells [7]. The same year, Mansoor *et al.* reported the isolation of three novel β-carboline indole alkaloids (92-94) from the MeOH extract of the leaves of *T. elegans* (Scheme 6) and these compounds were evaluated for their ability to modulate multidrug resistance in mouse lymphoma cell lines [41]. Moreover, this group was also reported the isolation a novel (95) and three known monoterpene indole alkaloids (96-98) from the methanol extract of *T. elegans* leaves. The isolated alkaloids were evaluated for their apoptosis induction activity in human hepatoma HuH-7 cells. The monoterpene indole alkaloids, 96 and 98, showed the most promising apoptosis induction activity in human hepatoma HuH-7 cells [42].

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**Scheme 7: Bioactive indole alkaloids (96-111) isolated from *Tabernaemontana* plants**

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**Tabernaemontana elegans Staph.**

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and 102, corroborating the nuclear morphology evaluation essay [43]. In 2011, Luo et al. reported the antimicrobial activity of selected medicinal plants traditionally used in Mozambique. From the results, the EA extract of T. elegans showed strong activity against Mycobacterium tuberculosis H37Rv at MIC value of 15.6 µg/mL and the prominent compounds were identified as indole alkaloids [44]. Moreover, the next year, Pallant, Cormarthy, and Steenkamp reported the isolation and identification the fraction responsible for the antimicrobial activity in T. elegans roots extract. It was found that the GC-MS analysis identified the indole alkaloids, dregamine (97) and voacangine (100), as major components which were shown good antibacterial activity including Gram-positive bacteria and Mycobacterium species, with MIC values in the range of 64–256 µg/mL [45].

**Tabernaemontana hystrix Steud.**

In 2010, Souza et al. reported the isolation of two new monoterpene alkaloids, ibogamine-7,8-dione (103) and 12-methoxyvobasoline (104) of the CHCl3 extract from the root bark of T. hystrix (Scheme 7). The structures of all compounds were elucidated on the basis of spectroscopic data analyses, mainly 1H- and 13C-NMR, including two-dimensional experiments [H, H-COSY, HMBC, and HMQC] [46].

**Tabernaemontana sessilifolia (Baker)**

In 2012, Girardot et al. reported the isolation of four vobasylé-boga bisindole (105-108) and one 2-acyl monomeric indole (109) alkaloids from the stem bark of T. sessilifolia (Scheme 7). All isolated alkaloids were evaluated for antiplasmodial activity against the chloroquine-resistant strain Fcb1 of Plasmodium falciparum, and cytotoxicity against the human lung cell line MRC-5 and the rat skeletal muscle cell line L-6. The alkaloid, namely, 3'-oxotabernaelegantine A (106) exhibited antiplasmodial activity (IC50 = 4.4 µM) associated with non-significant cytotoxicity (selectivity index of 48). Tabernaemontane genine B (101) displayed the highest cytotoxicity with IC50 values of 0.47 and 0.42 µM on MRC-5 and L-6 cells, respectively [47].

**Tabernaemontana sphaerocarpa Blume**

In 2009, Zaina et al. reported the isolation of biscarpamontamine A (110), possessing an aspidosperma-iboga-type skeleton, and biscarpamontamine B (111), having an aspidosperma-aspidosperma-type skeleton, from the stems of T. sphaerocarpa (Scheme 7). Bis-indole alkaloid 111 showed potent cytotoxicity against various human cancer cell lines, including human blood premyelocytic leukemia (HL60), multiple myeloma (RPMI8226), non-small cell lung carcinoma (NCI-H226), human colon cancer (HTC16), and human breast adenocarcinoma (MCF-7) cells with IC50 value in the range of 0.5-1.9 µM [48].

**CONCLUSION**

Recent studies have provided evidence that natural-derived bioactive monoterpene indole alkaloids play a vital role in human health and nutrition as well as disease prevention. They offer a diverse range of structurally distinctive bioactive molecules that have been used as a major source of innovative and effective therapeutic agents. Furthermore, the biological screening of active monoterpene indole alkaloids, using a wide variety of scientific tools and the intensive collaboration of experts in diverse scientific disciplines will become research hotspot, providing new and essential health-care opportunities.

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**REFERENCES**

26. Ngor E, Chong KW, Thomas NF, Lim SH, Low YY, Kam TS. Ibobgan,


