Article type : Special Issue Article



REVIEW ARTICLE

Title: The protective effects of Kava (Piper Methysticum) constituents in cancers: a systematic review

Running title: Kava, the pacific drug on the anti-cancer radar

Key words: kava, piper methysticum, cancer, flavokavains, oral cancer

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Abstract

This is the author manuscript accepted for publication and has undergone full peer review but has not been through the copyediting, typesetting, pagination and proofreading process, which may lead to differences between this version and the <u>Version of Record</u>. Please cite this article as <u>doi: 10.1111/JOP.12900</u>

Background: Kava is a beverage made from the ground roots of the plant Piper Methysticum and has long-held a significant place within Pacific island communities. Active compounds extracted from kava, and secondary metabolites, include kavalactones, chalcones, cinnamic acid derivatives and flavanones. It is thought that components of kava may exert an antiproliferative effect through cell cycle arrest and promotion of apoptosis.

Methods: We conducted a systematic review to summarize available evidence of the anticancer effects of kava components, and investigate their potential use for oral squamous cell carcinoma (OSCC) treatment. Eligible studies were identified through a comprehensive search of OVID EMBASE, OVID MEDLINE and Web of Science, as at April 2018.

Results: Of 39 papers that met the inclusion criteria, 32 included in-vitro models and 13 included animal studies. A total of 26 different cancers were assessed with 32 studies solely assessing epithelial cancers, 6 mesenchymal cancers, and 1 study including both. There was only one report assessing an OSCC cell line. Anti-proliferative properties were demonstrated in 32 out of 39 papers. The most researched constituent of kava was flavokavain B followed by flavokavain A. Both were associated with increased expression of pro-apoptotic proteins and decreased expression of anti-apoptotic proteins. Further, they were associated with a dose-dependent reduction of angiogenesis.

Conclusion: There was heterogeneity of study models and methods of investigation across the studies identified. Components of kava appear to present an area of interest with chemo-therapeutic potential in cancer prevention and treatment, particularly for epithelial neoplasms. To date, there is a paucity of literature of the utility of kava components in the prevention and treatment of oral squamous cell carcinoma.

Introduction

Kava is a beverage made from the ground roots of the plant *Piper Methysticum* (Fig. 1), directly translated as "intoxicating pepper". The word 'Kava' originates from Tongan and Marquesan and means "bitter".¹ Common names of Kava include: Kava; Kava-kava; Avaava; Antares; Ava; Ava pepper; Ava root; Awa; Fijian kava; Gea; Gi; Grog; Kao; Kava kava rhizome; Kava root; Kavapiper; Kavapyrones; Kavarod; Kavekave; Kawa; Kawa kawa; Kawa pepper; Kawa Pfeffer; Kew; Macropiper latifolium; Malohu; Maluk; Maori kava; Meruk; Milik; Pepe kava; Rhizoma piperis methystici; Sakaua; Sakau; Tonga; Yagona; Yangona; and Yongona.

Kava has long-held a significant place within Pacific island communities, being consumed for more than 2,000 years by individuals of Polynesian, Micronesian and Melanesian descent.²

The kava beverage is commonly drunk for social, ceremonial and medicinal purposes, acting as a muscle relaxant and inducing sleepiness and relaxation.³ Kava plays a valuable role as an item for exchange within political, religious and economic spheres. Traditional regulations existed on kava usage regarding gender, social class, religion and age.⁴ The use and cultivation of Kava has extended from traditional to recreational use, particularly in the last couple of decades.^{4,5} South Pacific islanders can consume Kava on a daily basis. Pacific island economies are reliant upon Kava both as a means of traditional subsistence and as a contemporary cash-crop and valuable export commodity.⁴

Recreational Kava usage has reached regions of Australia and New Zealand partly due to migration of Pacific Islander communities to these areas.⁶ Kava was also introduced as an alcohol alternative to Australian Aboriginal communities residing in Arnhem Island in 1982 in an attempt to reduce alcohol-related harm in the community.^{7,8} The import and purchase of Kava in Australia was unregulated until 6th of April 1998. The National Code of Kava Management's standard 2.6.3, ⁹ enabled individual states and territories to implement more restrictive measures. Revelation of hepatotoxicity risk followed safety assessments of adverse changes associated with excessive consumption.⁹ WHO's safety review of Kava has held concerns over the introduction of non-traditional preparations of kava into "exposure regions" such as Arnhem Island, in the Northern Territory where excessive consumption occurs. Currently, import, advertising and sale of Kava in Australia and New Zealand is strictly controlled under the Customs (Prohibited Imports) Regulations Act.¹⁰ Commercial importations are no longer allowed, with the exception of medical or scientific purposes and trafficable quantities for those without a license or permit is 2 kg.¹⁰

Kava is traditionally prepared using the rhizome of the plant, which can be used fresh or dried. If using fresh preparations, the roots are macerated and the resultant juices mixed with other solvents such as coconut milk or water. The more common style of consumption is obtained by adding water to finely ground kava roots followed by filtration to obtain the desired beverage.¹¹ Fresh preparation produces a stronger beverage than when prepared from dried starting material.¹²

Active compounds extracted from kava include 18 kavalactones, three chalcones, cinnamic acid derivatives and flavanones.¹³ Recent studies have identified 30 secondary metabolites, expanding to 19 kavalactones, 3 dihydrochalcones and 8 minor components.¹⁴ Ninety-six per

cent of the organic extract derived from kava rootstock consists of six major kavalactones, methysticin, dihydromethysticin, kavain, dihydrokavain, demethoxyyangonin, and yangonin (Fig. 2).¹⁴ Traditional kava preparations contain 0.3-20% kavalactones. Kavalactone and chalcone concentrations are increased when extracts are unfiltered.¹³ Commercialised kava preparations extracted by organic solvents demonstrate a similar compositional ratio, excluding greater kavain and dihydrokavain content.¹⁵

Kava is commonly known for its anxiolytic relaxant effects and these are achieved by kavalactone ligation to the central nervous system (CNS) GABA receptors.¹⁶ Modulation of GABA activity is mediated via alteration of lipid membrane structure, sodium channel function, MAO-B inhibition, and sodium and dopamine reuptake inhibition within the brain.¹⁷ This disruption of the GABA cerebellar function induces impaired movement coordination and visual attention that accompanies Kava inebriation. Further side effects associated with excessive consumption are dryness and skin ulceration.¹⁶

In addition to its psychotropic effects, recent evidence has shown that some of the Kava components exert anti-cancer effects. There is growing interest about their potential impact on malignant cell proliferation, signalling, resistance to apoptosis, evasion of growth suppressors and angiogenesis. However, the potential anti-proliferative effects on oral squamous cell carcinoma (OSCC) has yet to be investigated. This systematic review aims to present the current literature surrounding Kava and its potential anti-cancer effects with a specific focus on OSCC.

Objectives

This qualitative systematic review aims to collate scientific evidence that constituents of Kava have anti-cancer properties and may be potential chemotherapeutic agents. The specific questions addressed in this systematic review were:

- 1) Do the constituents of Kava have anti-proliferative effects on cancer cells in vitro?
- 2) Do the constituents of Kava have anti-proliferative effects on tumours in animal studies?
- 3) Do the constituents of Kava have cancer prevention properties?
- 4) What known anti-cancer pathways do the constituents of Kava interact with?
- 5) Which Kava constituents have the greatest anti-cancer potential?

Methods

The Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA Statement) was used for the data collection and sorting process. A table based on the PRISMA flow diagram is included (Fig. 3).

Literature search

On April 9, 2018, a comprehensive search of electronic databases OVID EMBASE, OVID MEDLINE and Web of Science was performed. An identical search strategy was employed across all databases.

The aim of the search was to find *in vivo* and *in vitro* experimental studies, allowing a summative evaluation of the preventive and treatment effects of Kava on cancer.

Search queries involved keyword searches for all cancer related terms and various Kava names including: anticancer OR anti-cancer OR anti-proliferative OR antiproliferative OR cancer OR carcinoma OR tumorigenesis OR neoplasm OR malignan* OR metastas* AND yaqona OR sakau OR rauschpfeffer OR kava OR kawa OR piper methysticum OR tudei OR piper wichmannii.

A second keyword search query was also performed for all cancer related terms and various active Kava constituents: anticancer OR anti-cancer OR anti-proliferative OR antiproliferative OR cancer OR carcinoma OR tumorigenesis OR neoplasm OR malignan* OR metastas* AND chalcones OR flavokawain OR flavokavain OR kavalactones OR desmethoxyyangonin OR 5,6-dehydroka* OR dihydromethysticin OR dihydrokavain OR hydroxykavain OR kavain OR kawain OR methysticin OR hydroxykavain OR hydroxykawain.

Selection criteria

No limits were placed on the search, and all languages were accepted. Papers that studied interactions between major kava constituents and cancer cells, *in vitro* or in animal models, were included. Technical reports were accepted. Studies using Kava constituents originating from other plant species were also included. No restriction was placed on what types of cancers were involved. Cell markers, cell proliferation/apoptosis and tumour sizes (*in vivo*) were common biomarkers. No restrictions were placed on the study design employed.

Studies were excluded if they were irrelevant to the cancer and Kava association. Papers studying the interaction of Kava or its constituents in non-cancer areas such as anxiety, hepatotoxicity or structure activity relationships were disregarded. However, the association between cancer and inflammation led to inclusion of papers discussing inflammatory effects of kava constituents if there was a concurrent focus on cancer. Studies of synthesised analogues of Kava constituents were not included, nor were studies solely focused on whole Kava extracts or fractions due to the large number of unknown variables that would be introduced. Case reports, letters, conference abstracts, reviews, epidemiological studies and retracted studies were excluded.

Data collection and sorting

The literature search identified a total of 2952 papers. Search results were imported into Endnote X8, compiled, and duplicates removed by the software, and then further scanned individually to remove manually undetected duplicates by two blind reviewers. A total of 1834 unique articles were found.

Utilizing the PRISMA protocol, 2 independent and blinded reviewers were enlisted to sort the relevance of articles by title.

Title screening

Out of the 1834 papers, 107 unique articles were selected with an inter-rater reliability (IRR) of 98.26%. If there was no accordance on one paper, it was evaluated by a third reviewer. A final total of 94 articles was unanimously selected for abstract screening.

Abstract screening

Of the 94 articles screened, a combined total of 55 articles were accepted (IRR: 92.55%). Disagreements were due to each reviewer's subjectivity. Discussion of the 7 differences resulted in a consensus of 3 articles being accepted. A final total of 51 articles was unanimously selected for full-text screening.

Full-text screening

4 reviewers screened the articles for data extraction to be included in the systematic review. Contentious articles were discussed by all reviewers. Unanimous agreement was required for the articles to be included in the final result.

Of the 51 articles screened, a combined total of 39 articles were accepted by the reviewers. Data extracted from each article was recorded on the data collection table displayed as table 1. The following information was collected from each article: active kava molecules, model systems employed consisting of *in vitro* and *in vivo* models, experimental model as well as major findings incorporating anti-cancer mechanisms identified.

Risk of bias across studies

As with other bodies of scientific evidence, the potential effect of publication bias, favouring reporting of positive outcomes, cannot be excluded. Likewise, it is not known whether the authors reported only their most favourable results. Therefore, included studies underwent quality assessment according to QUIPS guidelines. Averaged across each risk of bias domain, we evaluated 85.9% percent of articles as having a low risk of bias, 11.54% of articles were identified to have a moderate risk of bias, and 2.56% were at a high risk of bias (Appendix 1).

Results

The initial search strategy identified 2952 potentially relevant citations of which, 1118 duplicates were removed. Following screening by title and abstract, 51 publications were included for full-text screening by 4 independent reviewers against the eligibility criteria. 12 publications were excluded and the remaining 39 were included for data extraction in the systematic review. Rationale for exclusion is presented in Appendix 2.

Of the 39 included studies, Flavokavain B (FKB) was the most studied constituent (21 articles); followed by Flavokavain A (FKA) (12 articles); dihydromethysticin (DHM) (7 articles) and desmethoxyyangonin (DMY) (4 articles); dihydrokavain and kavain (5 articles each); Flavokavain C (FKC) (4 studies); methysticin (4 articles); yangonin (3 articles); and yangonindimers (1 article). A single study was also conducted on the recently isolated yangonindimers. A complete list of the included studies are presented in Table 1.

Thirteen articles included animal models; these included tumour-inoculated and dietary NNK-induced mice models (A/J, TRAMP, C57BL/6, UPII-SV40T)^{19 20,30,31,34,37-39,43}, *in situ* human cancer tissue xenografted into SCID mice and nude mice and the Balb/C strain ^{15,32,49,50}; and the zebrafish model.⁴⁵ Of the 13 articles conducting animal studies, 7 also studied the effects of kava constituents on cancer cells *in vitro*.^{15,19,20,31,32,49,50} Overall, 32 papers studied the effects of Kava constituents *in vitro*. Multiple cancer cell lines were studied, each representing diverse cancer types: breast, colorectal and lung cancer were commonly studied, while oral cancer was only studied in one publication.²⁷

Flavokavain B (FKB)

The most heavily researched constituent of Kava was found to be FKB. A total of 21 studies comprising 15 in vitro (including 2 in situ; 1 ex vivo), 1 in vivo, and 5 that assessed both models, examined the anti-cancer effects of FKB. Across these studies, there was unanimous agreement that FKB had anti-proliferative ability with the majority of papers reporting induction of G2/M arrest.^{14,18,22,23,25,27,29,32,49,50,52,53} One paper suggested FKB induced G0/G1 arrest.35 Additionally, general consensus that FKB there induced was а apoptosis.14,18,20,22,23,25,27,29,38-40 Proposed mechanisms included increased expression of proapoptotic proteins. such as PUMA, Bim and Bax expression ^{28,29,35,36,38}; decreased expression of anti-apoptotic proteins, such as survivin and XIAP^{33,35,36}, ROS production^{14,31,34}; or increase in caspase 3, 7, 8, and or 9.28,36 Three studies suggested apoptosis may have been induced via PARP cleavage ^{27,29,33}, although interestingly a single study reported that apoptosis was induced in the absence of PARP cleavage. ³⁴ A single study, found that FKB did not induce apoptosis.²⁰ Other proposed anti-cancer effects of FKB included prevention of metastasis^{19,30,32}, anti-angiogenic effects ^{27,30,32}, as well as regulation of immune and inflammatory functions.19

Flavokavain A (FKA)

A total of 12 studies comprising of 6 *in vitro* (including 1 *ex vivo*), 2 *in vivo*, and 4 that assessed both models examined the anti-cancer properties of FKA. FKA was found to induce apoptosis^{18,21,41-43} and inhibit proliferation.^{18,22,40-42} Proliferation inhibition was reported mainly through G2/M ^{21,42,43}, and G1 cell cycle arrest ²¹, as well as c-myc inhibition.^{18,40} Reported pathways inducing apoptosis included caspase activation^{40,41}, increase in pro-apoptotic protein expression such as Bim, Bax, DR5 and p27 ^{22,42}, decrease in anti-apoptotic protein expression namely survivin and XIAP^{22,40}, and various mitochondrial membrane changes. ⁴¹ Liu et al. (2017) found evidence of synergism in apoptosis induction between FKA and Yangonin.⁵⁰ Other anti-cancer findings of FKA included the prevention of metastasis, inhibition of angiogenesis, reduction in inflammation, and enhanced immune function.^{18,22,41} Solely, Johnson et al. (2011) concluded that FKA did not have significant anti-proliferative effects or apoptosis-inducing capabilities.²⁰

Flavokavain C (FKC)

A total of 4 studies, 3 *in vitro* and 1 that assessed both *in vitro* and *in vivo*, examined the anti-cancer properties of FKC. There was a consistent suggestion that FKC inhibited cell

proliferation in various cancer cell lines via G1 and G2/M arrest due to p21, p27, p53 upregulation.^{40,44} Furthermore, FKC *in vitro* induced apoptosis via down regulation of apoptosis inhibitors XIAP, c-IAP1, c-IAP2, upregulation of pro-apoptotic signals, such as CHOP and GADD153, increasing ROS, and decreasing superoxide dismutase.^{40,44,45} *In vivo*, FKC did not show anti-proliferative activity when administered alone to A/J mice.²⁰

Dihydromethysticin (DHM)

A total of 7 studies including 3 *in vitro*⁴⁶⁻⁴⁸ and 4 *in vivo*²³⁻²⁶, examined the anti-cancer properties of DHM. All of the studies demonstrated that DHM had chemopreventative effects, with one study suggesting induction of apoptosis by G0/G1 arrest.⁴⁷ The remaining *in vitro* studies reported weaker anti-cancer effects of DHM.^{46,48}

Methysticin

A total of 4 studies, 3 *in vitro* and 1 *in vivo*, examined the anti-cancer properties of methysticin. Methysticin has shown inhibition of NF-kB, a protein involved in cell survival, in A549 human lung adenocarcinoma, and on K562 chronic myelogenous leukaemia cell lines.^{46,48} Methysticin resulted in downregulation of androgen receptor genes, but not actual androgen receptor mRNA expression in prostate cancer cell lines.¹⁵ It was suggested methysticin's bioactivity is non-conformational,⁴⁶ and that its dioxymethylene group is essential to its activity.⁴⁸ All studies suggested methysticin may have promising anti-cancer activity, however the concentration tested and reported potency varied between studies.

Kavain

A total of 5 studies, 4 *in vitro* and 1 *in vivo*, examined the anti-cancer properties of kavain. Kavain was found to enhance growth inhibition of cancer cells, however with lesser potency than FKB and methysticin.^{15,48} Minimal to no anti-proliferative or apoptotic effects of Kavain was observed on three different bladder cancer cells in contrast to similar doses of flavokavain A, B and C. ⁴⁰ Finally, a single study found no effect of kavain on cell viability.⁴⁶

Dihydrokavain

A total of 5 studies; 3 *in vivo* and 2 *in vitro*; examined the anti-cancer properties of dihydrokavain. The studies unanimously agreed that dihydrokavain showed no significant anti-cancer effects.^{23-25,46,48}

Desmethoxyyangonin (DMY) (also known as 5,6-dehydrokavain)

A total of 4 studies, 3 *in vitro* and 1 *in vivo*, examined the anti-cancer properties of DMY. DMY demonstrated a highly selective anti-proliferative effect in ovarian and lung cancer.⁴⁹ In contrast, one study found that DMY concentrations of 5–20 μ g/ml did not have any effect on human squamous cell carcinoma of the cervix.²⁷ Finally, a study reported that DMY showed 100-300 times less NF- κ B inhibitory activity than methysticin.⁴⁸

Yangonin

A total of 3 studies, all *in vitro*, examined the anti-cancer properties of yangonin. Yangonin, was found to enhanced growth inhibition when combined with FKB, but demonstrated less potent growth inhibition independently.¹⁵ In comparison to FKB, yangonin had weaker bioactivity in both prostate cancer and K562 human chronic myelogenous leukaemia cell lines.^{15,46} Yangonin alone showed variable growth inhibition on murine cell lines as well as time and dose-dependent inhibition of human cell line growth.⁵⁰ When combined with docetaxel and FKA, yangonin synergistically induced autophagic cell death in bladder cancer cell lines.⁵⁰

Yangonindimers

One *in vitro* study assessed yangonindimers, involving the combination of recently isolated dimers of yangonin and desmethoxyyangonin. The study showed no significant anti-proliferative activity on several human tumour cell lines.⁵¹

Oral Squamous Cell Carcinoma

Hseu et al. (2012) was the only identified study to investigate components of Kava in OSCC. The study assessed the chemopreventive effect of FKB on two human tongue oral squamous carcinoma cell lines, HSC-3 and Cal-27. Major anti-proliferative measures assessed included cell viability determined by 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide (MTT) assay, cell cycle analysis by FACScan cytometer, and apoptosis by terminal deoxynucleotidyl transferase dUTP-fluorescein nick end-labeling (TUNEL) with fragmented DNA detection kit.³¹ It was found that FKB significantly inhibited growth of HSC-3 cells and Cal-27. However, the effect was more pronounced in HSC-3 cells, where it caused accumulation of cells in G2/M phase.³¹ Dose-dependent induction of apoptosis was also

observed.31

Discussion

Kava has been of interest for its established anxiolytic effects. ^{52,53} However, there is now a growing recognition of its potential anti-cancer effects. This systematic review evaluates the current literature on the anti-cancer properties of Kava constituents, their possible mechanisms of action, and areas that require further investigation. This review was focused on individual constituents as opposed to extracts. This has allowed identification of molecules of greatest chemotherapeutic significance and comparison between papers.

The gathered publications strongly supported that multiple constituents of Kava possess significant anti-proliferative and apoptotic properties that warrant further research. Anticancer properties were mainly assessed by cytotoxicity evaluation through cell viability tests, such as MTT assays, with protein analyses used to assess potential mechanisms involved in apoptosis. Numerous studies also investigated the impact on proliferation regarding the possible cell cycle inhibition by Kava including stage of cell cycle arrest. Methodological heterogeneity limited direct result comparison between studies, although this may be reflective of the novel interest in the anti-cancer properties of Kava.

Chemoprevention refers to the administration of an agent intended to reduce or delay the initiation of carcinogenesis.⁵⁴ Substances demonstrating anti-proliferative or apoptosis-inducing mechanisms, such as constituents of Kava, could have potential as chemo-preventive therapeutics. Anti-proliferative properties were demonstrated in 32 out of the studied 39 papers. Of these papers, 27 presented *in vitro* assessment and 10 *in vivo*, which were mainly conducted in mice, and a zebrafish model^{15,18-30}.

Fifteen papers reported cell cycle arrest by FKA, FKB, and or FKC, at G2/M^{14,21,27-29,31-38,42,43} or G1^{21,39}. Cells accumulated in the G2/M phase undergo apoptosis through the Bax-initiated mitochondrial pathway.^{33,38,42} Apoptosis was investigated in 24 out of 39 papers, 21 of which included *in vitro* components and 7 *in vivo* components. Apoptosis is characterised by key features, such as caspase activation, DNA fragmentation, and chromatin condensation.³¹ Many of kava's active compounds instigate apoptosis through DNA fragmentation and mitochondrial membrane permeability via upregulation of the mitochondrial apoptosis cascade and Fas-mediated apoptosis of Caspase-8.^{27,31,36} FKA, FKB and FKC induce apoptosis via the intrinsic pathway, associating with an increase in cytochrome c release leading to activation of Caspase 3, 7, 8 and 9.^{14,27-29,31,33,36,37,40,41,44,45} Caspase activation has

numerous effects, however the main mechanisms highlighted include PARP cleavage, increased ROS production, and consequent inhibition of PI3K/Akt phosphorylation in keratinocytes, reducing tumorigenesis.^{14,27-29,31,33,34,44}

In addition to increased caspase activation, FKA and FKB were associated with increased expression of pro-apoptotic proteins and decreased expression of anti-apoptotic proteins. Specifically, there was an increase in Puma, Bim, and Bax, activating BAK and BAX proteins responsible for the initiation of apoptosis at the mitochondrial outer membrane. ^{28,29,35,36,38,42} There were further reports of increased DR5 expression and subsequent caspase 8 activation, and upregulation of p21, p27 and p53, all which play a role in gene inhibition/cell cycle arrest and anti-proliferative pathways.^{21,27,35,43} FKA and FKB decreased the expression of anti-apoptotic proteins such as Bcl-2, BclX/L and Cdc2.^{14,27,36,42} Exposure to flavokavains proposedly lead to down regulation of the inhibitor of apoptosis (IAP) protein family, allowing for progression of apoptosis along with decreased inflammation.⁵⁵ BclX/L prevents the release of mitochondrial contents, such as cytochrome C, whilst Cdc2 triggers entry into the M phase of the cell cycle and suppresses the action of Wee1.⁴² ^{14,27,36} Wee1 typically slows admission into M phase to ensure adequate cell growth.^{21,27,42} Thus, Kava compounds have the capability to both downregulate anti-apoptotic and upregulate pro-apoptotic proteins with the potential to influence numerous apoptotic pathways.

A number of studies investigated angiogenesis-inhibitory properties and found Kava compounds successfully inhibited vessel formation *in vivo* and *in vitro*.^{18,22,30,32,41} Both FKA and FKB were shown to reduce angiogenesis in a dose-dependent manner.^{18,22,27,30,32,41} For instance, FKA was found to downregulate the androgen receptor and subsequently alter angiogenesis in bladder tumour tissues.²²

No studies reported the comparison of the anti-cancer capabilities of the constituents as a primary outcome. However, selected papers suggest that FKB may be the most potent constituent studied.¹⁵ One study concluded that kavain, DHK, DHM and DMY are approximately 100-300 times less active than methysticin⁴⁸ whilst a further study commented on the lesser toxicity of FKA in comparison to FKB.^{20,56}

Interestingly, a study undertaken by Johnson et al. (2011) on the effects of whole Kava extract in A/J mice found a statistically significant reduction in tumour multiplicity. However, they found that individual administration of FKA, FKB and FKC did not impact on adenoma multiplicity even at concentrations above that naturally occurring in Kava extracts.²⁰ Factors influencing these results could include variation in Kava extract, processing artefacts, strain of mice, tumour type and interactions of constituents and host

ligands. This reveals the complexity of *in vivo* models and the need for further studies to translate *in vitro* findings.

Within the included publications, a total of 26 different cancers were assessed across the 32 studies solely on epithelial malignancies, 6 on mesenchymal malignancies and one study including both.⁴⁹ A single publication analysed the effect of the Kava constituent FKB in ectodermal-epithelial cancers, particularly oral squamous cell carcinoma and melanoma.³¹ In contrast, the study of mesodermal-epithelial cancers and Kava constituents has been moderately established, particularly in colorectal and breast cancers.^{14,18,19,34,41} ^{42,44,45,49,51} Malignant epithelial lines studied may have potential mechanistic correlations in OSCC.

Mesenchymal cancer studies were the least investigated with primary focus on leukemia.^{46 29} Consideration can be given however, to their potential application in the context of epithelialmesenchymal transition in epithelial malignancy (EMT). The effect of Kava in sarcoma studies could reveal target pathways in invasive carcinomas, such as OSCC, demonstrating EMT with mesenchymal traits. For example, EMT has been implicated in metastasis and invasion by the PI3K/Akt pathways in OSCC.⁵⁷

Conclusion

This systematic review found that particular constituents of Kava consistently displayed anticancer effects *in vitro* and *in vivo*. The current body of evidence primarily features the induction of apoptosis and cell cycle arrest in epithelial malignancy. These promising findings highlight an avenue for further research of Kava constituents in the prevention and treatment of OSCC. There remains a paucity of literature surrounding the chemotherapeutic potential of Kava in the field of oral oncology.

Acknowledgement

The authors would like to thank Dr. Jim Berryman, Liaison Librarian, Brownless Biomedical Library, University Library, The University of Melbourne, for his guidance with the systematic review. The authors would like to thank the staff at the National Herbarium of Victoria (MEL), including the digitising officer Angharad Johnson and the manager of the herbarium specimen collection Pina Milne, who provided the image included in Figure 1.

Conflict of interest

The authors declare that they have no conflict of interest.

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Figure legend

Figure 1: *Piper Methysticum* pressed plant specimens dated back to 1886, from the collection of the National herbarium of Victoria collection. Images were captured with a Leaf Aptus-II 10 Digital Back camera. Reproduced with permission from the Royal Botanic Gardens Victoria.

PRISMA flow chart of the systematic review

PRISMA flow chart of

the systematic review

Figure 2: Structures of the major kavalactones occurring in kava rhizome; (below) ground kava root.

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Figure 3: PRISMA flow chart of the systematic review

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Ref.	Author,	Active molecule(s)	Model System	Experimental model	Major findings / mechanisms reported
	Year				
18	Abu et al	Flavokavain B	Human breast cancer	In vitro and ex vivo	- Induced apoptosis in both MCF-7 and
	2016		(MDA-MB231 MCE-7)	Cell culture MTT assay cell treatment BrdU	MDA-MB231: G2/M arrest was seen in
	2010			incorporation assay	MDA-MB231, C2/11 arcst was seen in
		\mathbf{O}		cell cycle analysis anneyin V/FITC assay wound	- Inhibited the in vitro migration and
		()		healing assay transwell migration/ Invasion	invasion in MDA-MB231 cells in a dose
				assay, HIWEC tube formation assay, as vivo rat	dependent menner
				assay, no vec tube formation assay, ex vivo fat	Inhibited angiogenesis: via suppressing the
				aortic ring assay, quantitative rear-time PCK,	- Initioned angiogenesis, via suppressing the
		<u> </u>		western blot analysis, proteome promer array	iormation of vessels in HUVEC cells and
		σ			within the rat aortic ring assay
19	Abu et al.,	Flavokavain A	Mouse breast cancer (4T1),	In vitro	- Dose dependant inhibition of proliferation
	2015a		Mouse lymphoma (YAC-1)	MTT assay, Wound healing assay, Transwell	of 4T1 cells
				migration/ invasion assay	- Anti metastatic effects in vitro
		<u> </u>			- Decreased volume and weight of tumors in
		0		In vivo	vivo
		0		TUNEL assay, tumor histopathology staining,	- Induced apoptosis in vivo
	_			immunophenotyping of splenocytes, splenocytes	- Suppresses proto oncogene C-myc
				cytotoxicity assay, serum detection of IL-2 and	- Enhanced T cell immunity via upregulation
				IFN-γ levels, Nitric oxide detection assay,	of interleukin 2 and interferon gamma,
				Quantitative real time PCR analysis, Western blot	resulting in an increase Th, Tc, NK cell
					populations
					- Anti inflammatory effect via reduction in
					COX-2 expression
					<u>r</u>

20	Abu et al.,	Flavokavain B	Mouse breast cancer (4T1)	In vitro	- Inhibited proliferation and induced
	2015b			MTT analysis, cell cycle analysis, wound healing	apoptosis of 4T1 cells
		<u> </u>		analysis, in vitro migration/ invasion assay,	- Inhibition of migration and invasion of 4T1
		\bigcirc		animal and diet, tumour inoculation and	cells
	-			treatment, terminal deoxynucelotidyl transferase	- Regulates several immune system markers
		<u> </u>		dUTP nick end labeling analysis of the tumours,	- Possesses antimetastatic abilities
		()		haematoxycilin and eosin histology staining, nitric	- Regulates inflammation and metastasis-
		6		oxide detection, lung, liver and spleen clonogenic	related genes and proteins
		0)		assay, immunophenotyping, serum detection,	
				bone marrow smearing, total red blood cell count,	
		2		real-time quantitative polymerase chain reaction,	
				western blot analysis, proteome profiler mouse	
		σ		angiogenesis, TUNEL analysis	
21	Abu et al.,	Flavokavain A	Human breast cancer (MCF-	In vitro	- Dose-dependently inhibits MCF-7 and
	2014		7, MDA-MB231)	MTT assay, BrdU assay, Annexin V analysis, cell	MDA-MB231 cell proliferation
		_		cycle analysis, JC-1 mitochondrial dye, AO/PI	- Cell-cycle accumulation at the G2/M phase
				dual staining, caspase 8/9 fluorometric assay,	in MDA-MB231
		\bigcirc		PCR, Western blot	- Dose dependently induced apoptosis in
					MDA-MB231 and MCF-7 cells through
	-			In vitro and ex vivo	mediation of caspase 8 and 9, and regulation
				Scratch assay, transwell migration/invasion assay,	of several apoptosis genes and proteins
				HUVEC tube formation assay, ex vivo rat aortic	- Induced changes in mitochondrial
				ring assay, quantitative PCR, Western blot	membrane potential. Higher doses reduces
					membrane polarisation
		~			- Inhibited motility and invasiveness of

					MDA-MB231 cells in vitro
					- Dose-dependently reduced angiogenesis
22	An et al.,	Flavokavain B	Human lung cancer (H460),	In vitro	- Inhibited cell proliferation
	2012		primary mouse embryo	MTT assay, cell morphology observation and	- Induced G2-M cell cycle arrest
			fibroblasts (MEF) deficient	DAPI staining,	- Induced apoptosis; through Bax initiated
		$\overline{\mathbf{a}}$	for Bax (double knockout)	fluorescence-activated cell sorting analysis of cell	mitochondrial pathway, significantly
		U		cycle distribution, Western blot analysis, plasmid	decreased the levels of survivin and X-linked
		S		transfection	inhibitor of apoptosis (XIAP), cytochrome c
					release and activated the cleavage of PARP,
					caspase-7, and caspase-9
					- Disruption of Bax almost completely
		T			impaired effect of FKB-induced growth
	_				inhibition (MEFs)
					- Activated the stress-responsive mitogen
					activated protein kinases (MAPKs)
					signalling
23	Chang et	Flavokavain B	Human gastric carcinoma	In vitro	- FKB is potently cytotoxic to human gastric
	al., 2017	9	(AGS, NCI-N87, Kato-III,	Cell culture, drug treatment, MTT assay, colony	cancer cells, mildly toxic towards normal
	_		TSGH9201), primary mouse	formation assay, measurement of ROS generation,	(Hs738) cells and primary mouse
			hepatocytes, normal	cell-cycle analysis, western blot analysis, GFP-	hepatocytes; induced AGS cell death
			stomach and intestinal cells	LC3 plasmid transfection and GFP-LC3 dot	characterized by autophagy as evidenced by
			(Hs738), athymic nude mice	formation	increased LC3-II accumulation, GFP-LC3
	•	Z	(BALB/c-nu) with AGS		puncta and acidic vesicular organelles
			tumour cell inoculation	In vivo	(AVOs) formation, without resulting
				Acridine orange staining, transfection of shRNA	procaspase-3/ PARP cleavage

	-	IUSCript		targeting LC3, Bax transfection, apoptotic DNA fragmentation, tumour cell inoculation animals, histopathological analyses of xenografted tumour, western blotting of xenografted tumours, animal survival study	 FKB induced apoptosis via ROS generation and ROS inhibition; indicated by ROS- mediated autophagy in AGS cells FKB induces G2/ M arrest and alters cell- cycle proteins; ROS-JNK signaling FKB inhibits apoptotic Bax expression, and Bax-transfected AGS cells exhibit both apoptosis and autophagy; thus, FKB- inactivated Bax
24	Dai et al., 2015		Human osteosarcoma (MG- 63)	In vitro MTT assay, computer-assisted phase, contrast microscopy, annexin V-FITC assay, flow cytometry, mitochondrial membrane potential measurement, western blot analysis	 Dose dependently inhibited growth of MG- 63 cells via chromatin condensation. Induced both cytotoxic and cytostatic effects resulting in marked vacuolization processes and ultimately cell death The percentage of cells in G2/M phase decreased considerably with increase in DHM dose whilst fraction of G0/G1 cells increased significantly with increasing concentration of DHM The number of cells with depolarized mitochondria increased with DHM dose
25	Eskander et al., 2012	Flavokavain B	Human uterine leimyosarcoma (SK-LMS- 1), human endometrial	In vitro MTT assay, Fluorescence activated cell sorting analysis of apoptosis, Fluorescence activated cell	- Selective inhibition of growth, greater effect in SK-LMS-1 and ECC-1 than in T- HESC

		adenocarcinoma (ECC-1),	sorting analysis of cell cycle, Western blot	- Induced G2/M arrest and apoptosis via
		human normal endometrial	analysis, Real-time reverse transcription-	upregulation of DR5, puma and bim
		fibroblasts (T-HESC)	polymerase chain reaction	expression, downregulation of apoptosis
	\mathbf{O}			inhibitor survivin
26	Folmer et Kavain, Flavokavain	Human chronic	In vitro	- Yangonin has weak bioactivity suggesting
	al., 2006 A, Flavokavain B,	myelogenous leukemia	Transient transfection, luciferase reporter gene	methoxy group of A ring hinders its activity
	Methysticin,	(K562), Human T cell	assay, Electrophoretic mobility shift assay	- compounds with aromatic rings solely
	Dihydromethysticin,	leukemia (Jurkat)	(EMSA), Western blot analysis, Protein Kinase	substituted with a methoxy group tend to
	Yangonin,		assays	lack anticancer activity.
	Dihydrokavain,			- FKA inhibits MAPKAP-K3 - could be
	Demethoxyyangonin			responsible for apoptotic mechanisms
	g			induced by Kava.
27	Hseu et al., Flavokavain B	Human oral squamous	In vitro	- Significantly reduced HSC-3, A-2058, Cal-
	2012	carcinoma (HSC-3), human	MTT colorimetric assay, Flow cytometric	27 cell survival in dose-dependent manner,
		melanoma (A-2058), oral	analysis, TUNEL assay, Fluorescent imaging of	minimal effect against A-549. Time- and
		adenosquamous carcinoma	mitochondria and endoplasmic reticulum,	dose-dependent for HSC-3
		(Cal-27), human lung	Immunofluorescence assay, Fluorescence	- Promotes growth inhibition by G2/M phase
		carcinoma (A-549)	microscopy for apoptosis, Western blot	arrest; inhibits cell cycle progression by
				reducing levels of cyclin A, cyclin B1, Cdc2,
				and Cdc25C (HSC-3)
				- Induces apoptosis (HSC-3); induces
				Apoptotic DNA Examplettion $(USC 2)$
				Apoptotic DNA Fragmentation (HSC-5),
				induces mitochondrial
	A			induces mitochondrial membrane permeability (HSC-3),
	A			induces mitochondrial membrane permeability (HSC-3), upregulates Mitochondrial Apoptotic

	anuscript			Cascades (HSC-3), activates Fas-Mediated Apoptosis through the activation of Caspase- 8 - Downregulates p38 MAPK and Upregulates ERK and JNK Proteins (critical in cell fate, Role in G2/M Arrest and Apoptosis, HSC-3) - Induces Intracellular ROS Generation (HSC-3) - Inhibits Phosphorylation of PI3K/Akt in HSC-3 (dysregulation of the PI3K/Akt signaling pathway leads to tumorigenesis in vitro and in vivo)
28	Jandial et al., 2017	Human breast adenocarcinoma (SKBR3), human breast cancer (MCF7/HER2)	In vitro MTT assay, Soft agar colony formation, Flow cytometric analysis of cell cycle distribution, Western blotting analysis, In vitro kinase assay, DAPI nuclear staining	 G2M arrest and apoptosis induction via downregulation of HER-2 gene that is known for suppressing proapoptotic proteins, increased expression of proapoptotic proteins Bim and BAX, decreased expression of anti apoptotic proteins Bcl₂, BclX/L, XIAP, survivin, dephosphorylation of Cdc25C, downregulation of Cdc2 hence Myt1 and Wee1 expression
29	Ji et al., Flavokavain B 2013	Human osteosarcoma (OS160, 143B , SaOS-2, MG-63, U2OS), Murine	In vitro MTT assay, Soft agar colony formation assay, DAPI staining for apoptotic cell nuclei, Caspase	 Inhibits proliferation of osteosarcoma cells Induces apoptosis (143B, SaOS-2) through increased expression of Fas, Puma and Bax,

			bone marrow cells (derived	activity assay, Fluorescence-activated cell sorting	while down-regulating the expression of Bcl-
			from Balb/c), Human normal	(FACS) analysis, Protein isolation and Western	2 and Survivin, increases Caspase 8, 9, 3/7
			small intestine epithelial	blot analysis, Zymogram assay, Motility and	activity (143B, SaOS-2)
		\mathbf{O}	cells (FHS)	invasion assay	- Suppressed in vitro motility (143B) and
	=				invasiveness (143B, SaOS-2); Inhibited
					MMP-2 and MMP-9 secretion (143B)
		\mathbf{O}			- Induces G2/M arrest (143B, SaOS-2 cells);
		6			Caused significant decrease in Cyclin B1,
		07			Cdc 25c and increase in p-Cdc2 in a time-
					dependent manner, increase in Myt1 not time
					dependent (143B)
					- No significant growth inhibitory effects on
		σ			murine bone marrow cells
					- Significant differences in cell viability for
		\geq			143B and FHS, both reducing with FKB
30	Johnson et	Kava extract	A/J mice with	In vivo	- Kava extract dose dependently reduced
	al., 2011	Flavokavain A,	dietary 4-	Tumour count by dissecting microscope, detection	lung adenoma multiplicity
		Flavokavain B,	(methylnitrosamino)-1-(3-	of serum enzymatic levels of alanine transaminase	- FKB significantly decreases adenoma
		Flavokavain C	pyridyl)-1-butanone (NNK)	(ALT), aspartate transaminase (AST), alkaline	multiplicity but at much higher concentration
	-		and benzo(a)pyrene (BaP)-	phosphatase (ALP), and total bilirubin, Western	than found in kava; FKA and FKC not
		1	induced lung tumorigenesis	Blot analysis,	significant effect
				immunohistochemistry staining; H&E staining	- No significant change to diet consumption,
					≤6% weight loss for highest kava
		Y			concentration, No significant change to liver
					weight and serum enzyme levels

)t			 No induction of apoptosis (no PARP cleavage) Dose dependent decrease in PCNA protein
14	Kuo et al., 2010	Human wild type and p53–/– colorectal carcinoma (HCT116), human lung carcinoma (A-549), mouse fibroblast (NIH3T3), human fibroblast (HFW)	In vitro Colony formation assay, Apoptosis assays, Confocal microscopy, Cytofluorimetric analysis of mitochondrial membrane potential, Measurement of intracellular calcium concentrations, Measurement of ROS, Cell cycle analysis, Western blot analysis, Real-time reverse transcription - polymerase chain reaction	 Concentration and time dependant cytotoxic effect Induction of G2/M accumulation, autophagy, and the intrinsic apoptosis pathway via the initiation of ROS-mediated apoptosis, GADD153 up-regulation resulting in ER stress response and downregulation of antiapoptotic Bcl-2 expression, induction of phosphorylation of p38 MAPK, and induction of cytochrome c release and Bak translocation resulting in mitochondria dependant apoptosis
31	Li et al., 2015	Human prostate adenocarcinoma (DU145, PC3, 22Rv1), prostate stromal cells (PrSCs), TRAMP mice (C57BL)	In vivo Western blot analysis, Plasmid and siRNA transfection, TUNEL assay In vitro MTT assay, NEDD8 conjugation assay, molecular docking, western blot analysis	 Induced G2-M arrest through activation of Myt1, Wee1, and CDK1 Induced G1 arrest via decreasing cyclin- dependent kinase-2 (CDK2) kinase activity, increasing p21/WAF1 and p27/ KIP1 expression, reducing CDK1-inhibitory kinase expression Degraded SKP2; a protein associated with tumor development, via induction of proteasome dependent degradation

15	Li et al.,	Kavain, 5',6'-	Human prostate cancer	In vitro	- Inhibits growth of AR-expressing PCa
	2012	dehydrokavain,	(LNCaP, LAPC-4, 22Rv1,	MTT Assay, Western Blot analysis, Quantitative	cells; Kava root extract more potent than
		yangonin,	PC-3, DU145, C4-2B)	RT-PCR, Transfection, promoter activity and	kavalactones (K, Y, 5'6-DHK, M) but less
		methysticin,	Human prostate normal	luciferase assay, Chromatin immunoprecipitation	potent than FKB
	=	Flavokavain B	(WPMY-1)		- Decreases expression of AR target genes
				In situ	PSA and TMPRSS2 through acceleration of
		Kava root extract	Tumour xenograft in SCID	Serum prostate-specific antigen, Caliper	AR protein degradation (Kava root extract
		6	mice	measurement of tumor size	and kavalactones)
		07	(GM0308, RC0309)	Immunohistochemistry staining	- FKB down-regulates expression of AR and
					its target genes (PSA and TMPRSS2)
					- Kavalactones (K, Y, 5'6-DHK, M) and
					FKB combination results in enhanced
		σ			inhibitory effect on growth of C4-2B cells
					and expression of AR protein
		>			- Kava extract and FKB decrease growth of
					patient-derived PCa xenografts in SCID
					mice, AR expression in tumor tissues, serum
		\overline{O}			PSA levels
32	Lin et al.,	Flavokavain B,	Human squamous carcinoma	In vitro	- FKB induced cell death (viability or
	2012	Desmethoxyyangonin	from glandular cancer of	Terminal deoxynucleotidyl transferase-meditated	growth) in a dose- and time-dependent
			cervix (KB), human	dUTP nick end-labeling assay for DNA apoptotic	manner
			gingival fibroblast (HGF)	fragmentation, Flow cytometric analysis (incl.	- DMY concentrations of 5–20 $\mu g/ml$ did not
				Cellular DNA content, Mitochondrial membrane	affect KB cells at 24h
			Tumour xenograft in nude	potential), Fluorescence microscopy/flow	- FKB cytotoxic to HGF above 30µg/ml at
		7	mice	cytometry measuring ROS generation, Western	24h

				Blotting, Zymography (MMP-9 activity),	- FKB induced apoptotic DNA
					fragmentation, release of cytochrome c,
				In situ	activation of caspase-3 and -9 and cleavage
		\mathbf{O}		Apoptosis detection by TUNEL with the Klenow	of PARP, activation of the Fas-mediated
	-			DNA fragmentation detection	apoptosis pathway by FKB results in
					activation of caspase-8 and cleavage of Bid,
		()			induces dysregulation of Bcl-2 and Bax
		000			protein, induced ROS generation and
					mitochondrial dysfunction
					- Sub-G1 accumulation and G2/M arrest in
					FKB-treated KB cells; dose- and time-
					dependent reductions in mitotic cyclins A
		M			and B1, mitotic-cyclin-dependent kinase
					Cdc2 and mitotic phosphatase Cdc25C
		>			expression, increases expression of
					p21/WAF1, Wee1 and p53
					- FKB dose-dependently affects metastasis-
					related protein expression; reduction of
		0			MMP-9 and u-PA expression, upregulation
		L			expression of specific endogenous inhibitors
					TIMP-1 and PAI-1
					- FKB In vivo inhibition of KB xenograft
					growth (tumour volume); apoptotic DNA
	•	A			fragmentation in xenograft tumors
33	Liu et al.,	Flavokavain A,	Human bladder cancer (T24,	In vitro	- Yangonin and 5' 6'-dehydrokavain are

	2017	yangonin, 5' 6'-	RT4, UMUC3, HT1376,	MTT assay, colony formation assay,	potent inducers of autophagic cell death in
		dehydrokavain	HT1197), mouse embryonic	western blot analysis, 7-Methyl-guanosine cap	bladder cancer cells
			fibroblasts (MEFs)	binding assay, stable transfection, fluorescence	- Yangonin induces autophagy via increased
		\mathbf{O}		microscopy, electron microscopy	expression of beclin and ATG5.
	=				- Yangonin reduces the viability of bladder
					cancer cell lines, and acts synergistically
		\mathbf{O}			with apoptosis-inducing agents such as
		6			docetaxel and flavokavain A; growth
		07			inhibitory effects of yangonin were
					attenuated in TSC1 or LKB1 knockout
					mouse embryonic fibroblasts, suggesting that
					TSC1 and LKB1 expression may contribute
		g			to optimal growth inhibition by yangonin
34	Liu et al.,	Flavokavain A	Transgenic mouse model	In vivo	- Concentration and time dependant anti
	2013		(UPII-SV40T)	Immunohistochemistry, DeadEnd colorimetric	tumor growth effect in vivo
		_		TUNEL assay, Western blotting	- Induced apoptosis via upregulation of
					proapoptotic protein DR5, p27 and TUNEL-
		0			positive apoptotic cells, downregulation of
					antiapoptotic proteins Ki67, survivin, and
	-				XIAP, downregulation of androgen receptor
		1			thus altering angiogenesis in bladder tumor
					tissues
					- Selective, no effect on organ:body weight
		Y			change, weight loss, or food / water
					consumption
34	Liu et al., 2013	Flavokavain A	Transgenic mouse model (UPII-SV40T)	In vivo Immunohistochemistry, DeadEnd colorimetric TUNEL assay, Western blotting	to optimal growth inhibition by yangonin - Concentration and time dependant anti tumor growth effect in vivo - Induced apoptosis via upregulation of proapoptotic protein DR5, p27 and TUNEL positive apoptotic cells, downregulation of antiapoptotic proteins Ki67, survivin, and XIAP, downregulation of androgen receptor thus altering angiogenesis in bladder tumor tissues - Selective, no effect on organ:body weight change, weight loss, or food / water consumption

35	Malami et	Crude extracts	Human colon cancer cells	In vitro	- UCK2 mRNA expression significantly
	al., 2017	FKB	(HT-29), human hepatic	Cell Viability study (MTT assay), RT-PCRof	downregulated in all HT-29 cells treated
		Alpinetin compounds	progenitor cells (HepaRG),	mRNA expression, Fluoresence imaging of 18S	with FKB and APN - consequent reduced
		(APN)	Monkey fibroblast-like	RNA expression, Cell Cycle analysis, DNA	18s rRNA in HT-29 cells.
	-		kidney cells (Vero)	Fragmentation analysis, Extraction of total	- G0/G1 phase inhibition by FKB
				protein, Western blot analysis	- FKB and APN induced p53 dependent
		\mathbf{O}			apoptosis.
		0			- MDM2 protein expression significantly
		07			downregulated with increased p53 protein
					expression in a time-dependant manner
					following FKB and APN treatment -
					essential in triggering cell cycle arrest and
		σ			apoptosis in HT-29 cells
36	Mustahil et	5,6-dehydrokavain,	Human lymphoblastoid	In vitro	- All crude extracts of the plant (5,6-DHK,
	al 2013	flavokavain B	cancer (T4 cells)	Cytotoxic assay, antimicrobial assay,	
	al., 2015	Havokavalli D,			FKB, pinostrobin and pinocembrin together
	al., 2013	pinostrobin,		DPPH radical scavenging activity assay	FKB, pinostrobin and pinocembrin together with Beta-sitosterol) demonstrated strong
	al., 2015	pinostrobin, pinocembrin, beta-		DPPH radical scavenging activity assay	FKB, pinostrobin and pinocembrin together with Beta-sitosterol) demonstrated strong cytotoxicity against CEMss (T4) cancer cells
	al., 2013	pinostrobin, pinocembrin, beta- sitosterol		DPPH radical scavenging activity assay	 FKB, pinostrobin and pinocembrin together with Beta-sitosterol) demonstrated strong cytotoxicity against CEMss (T4) cancer cells - FKB was the most cytotoxic isolate; most
	al., 2013	pinostrobin, pinocembrin, beta- sitosterol		DPPH radical scavenging activity assay	 FKB, pinostrobin and pinocembrin together with Beta-sitosterol) demonstrated strong cytotoxicity against CEMss (T4) cancer cells - FKB was the most cytotoxic isolate; most crude extracts and isolated compounds
	al., 2013	pinostrobin, pinocembrin, beta- sitosterol		DPPH radical scavenging activity assay	 FKB, pinostrobin and pinocembrin together with Beta-sitosterol) demonstrated strong cytotoxicity against CEMss (T4) cancer cells FKB was the most cytotoxic isolate; most crude extracts and isolated compounds showed weak activity in antimicrobial and
	al., 2013	pinostrobin, pinocembrin, beta- sitosterol		DPPH radical scavenging activity assay	 FKB, pinostrobin and pinocembrin together with Beta-sitosterol) demonstrated strong cytotoxicity against CEMss (T4) cancer cells FKB was the most cytotoxic isolate; most crude extracts and isolated compounds showed weak activity in antimicrobial and diphenylpicryllhydrazyl (DPPH) radical
	al., 2013	pinostrobin, pinocembrin, beta- sitosterol		DPPH radical scavenging activity assay	 FKB, pinostrobin and pinocembrin together with Beta-sitosterol) demonstrated strong cytotoxicity against CEMss (T4) cancer cells FKB was the most cytotoxic isolate; most crude extracts and isolated compounds showed weak activity in antimicrobial and diphenylpicryllhydrazyl (DPPH) radical scavenging activity tests
37	Narayanapi	pinostrobin, pinocembrin, beta- sitosterol Dihydromethysticin	A/J mice with dietary	DPPH radical scavenging activity assay In vivo	 FKB, pinostrobin and pinocembrin together with Beta-sitosterol) demonstrated strong cytotoxicity against CEMss (T4) cancer cells FKB was the most cytotoxic isolate; most crude extracts and isolated compounds showed weak activity in antimicrobial and diphenylpicryllhydrazyl (DPPH) radical scavenging activity tests DHM and methysticin significantly
37	Narayanapi Ilai et al,	pinostrobin, pinocembrin, beta- sitosterol Dihydromethysticin Dihydrokavain	A/J mice with dietary 4-(methylnitrosamino)-1(3-	DPPH radical scavenging activity assay In vivo DNA isolation following Genomic-tip 100/G	 FKB, pinostrobin and pinocembrin together with Beta-sitosterol) demonstrated strong cytotoxicity against CEMss (T4) cancer cells FKB was the most cytotoxic isolate; most crude extracts and isolated compounds showed weak activity in antimicrobial and diphenylpicryllhydrazyl (DPPH) radical scavenging activity tests DHM and methysticin significantly reduced NNK-induced O 6-mG while other

		, kavain, methysticin		lung tissues by LC-ESI-MS/MS, H&E staining	methylenedioxy functional group highlighted, minimum chemopreventive dose of DHM 0.01-0.1mg/g dietary - DHM significantly reduced NNK-induced adenoma multiplicity
38	Narayanapi llai et al, 2016a	Dihydromethysticin Dihydrokavain	Female mice with NNK induced lung adenoma (C57BL/6)	In vivo Quantification of O6-mG adduct in lung tissues, Urinary NNAL-O-Gluc and free NNAL quantificatoin, western blotting analysis, quantitative reverse transcription polymerase chain reaction	 DHM presented chemopreventive effects; DHM reduced the level of NNK and NNAL induced DNA damage in lung tissue but not liver tissue, and dose dependently reduced carcinogen (O6-mg) levels Suggests chemopreventive effect independent of Ahr pathway DHK had no effect
39	Narayanapi llai et al., 2016b	Dihydromethystycin, Dihydrokavain, Kava extract	A/J mice with dietary 4-(methylnitrosamino)-1(3- pyridyl)-1-butanone(NNK)	In vivo CYP2A5 Enzymatic Assays, Urine Processing to Convert NNAL-gluc into NNAL and Liquid Chromatography with Tandem Mass Spectrometry Quantification, Direct LC-MS/MS Detection and Quantification of NNAL-O-gluc, NNK, and NNAL in Urine and Serum Samples, NNAL Glucuronidation Activity of Mouse Lung and Liver Microsomes	 DHM and kava had no effect on NNAL formation from NNK Unlikely that the DHM primary mode of action is inhibition of CYP2A5-mediated NNAL bioactivation Higher amounts of NNAL-O-gluc detected in DHM-treated mice compared to control Enhanced NNAL glucuronidation activity in lung and liver tissues upon dietary DHM treatment DHK had no observable effect.

40	Phang et	Flavokavain C	Human colon	In vitro	- Induced reduction in cell proliferation
	al., 2017		adenocarcinoma (H-29),	Annexin V-FITC and PI assay, DNA	- Induced G1 and G2-M arrest via p21, p27,
		<u> </u>	human colorectal carcinoma	Fragmentation detection, Measurement of	p53 upregulation
		\bigcirc	(HCT 116)	mitochondrial membrane potential, Measurement	- Induced apoptosis via disruption of
	-	_		of intracellular ROS level, Superoxide dismutase	mitochondrial membrane potential activating
				inhibiton activity, Measurement of caspase -3,-8	caspases and PARP cleavage,
		\mathbf{O}		and -9, Cell cycle analysis, Western blot	downregulating apoptosis inhibitors XIAP,
		6			c-IAP1, c-IAP2, upregulating apoptotic
		0,			signal GADD153 via ER stress pathway,
					increased ROS, and decreased SOD
41	Phang et	Flavokavain C	Colon carcinoma cells (HCT	In vitro	- Reversing functional groups in FKC
	al., 2016	R	116),	Cytotoxic assay, Morphological assessment of	resulted in pronounced cytotoxic activity of
		(U	human colon	cell death by phase contrast and fluorescence	FKC in HCT 116 cells, cytotoxic and
			adenocarcinoma (HT-29),	microscopy, Plasma membrane, alteration	apoptotic activities of chalcones are
			human lung carcinoma	analysis, Assessment of changes in mitochondrial	dependent on its molecular structure.
		-	(A549), human cervical	membrane potential, Detection of DNA	- Gradual increase in cytosolic AIF and
			carcinoma (CaSki), human	fragmentation by TUNEL assay	Smac/DIABLO concentration with FKC
		0	breast adenocarcinoma	Assay for activation of Caspase-3/8/9, Cell Cycle	treatment. (AIF is involved in induction of
			(MCF7), non-cancerous	analysis using PI staining and flow cytometry,	caspase-independent chromatin condensation
	-		human colon fibroblast	Western blot analysis - analysis of apoptosis	and DNA fragmentation).
			(CCD-18Co)	related proteins in HCT116 following FKC	- Higher amounts of caspase-8 detected in
				treatment	comparison to active caspase-9 in HCT116
					cells with FKC treatment (dose-dependent).
		L			(Caspase 8 activation inhibits apoptosis
					triggered by death receptors.)

	-	Inuscript			 Increase in CHOP levels following FKC treatment. (CHOP is a key factor in ER stress-induced apoptosis via outer mitochondrial membrane permeabilization, caspase activation, and amplification of death signals) Suggested that an interplay between Akt signalling and MAPK apoptosis pathways and cell cycle arrest in HCT116 cells by FKC. FKC inhibited Akt activation, resulting in increased ERK1/2 phosphorylation. FKC activated p38 MAPK to some extent.
42	Pinner et al., 2016	Flavokavain A, Flavokavain B	Human hepatocellular carcinoma (HepG2)	In vitro RNA collection and real-time RT-PCR, Protein collection and Western blots, Total GSH assay, fluorescent viability	 FKA is considerably less toxic than FKB to HepG cells FKA and FKB increase total GSH levels FKA and FKB induce antioxidant and heat shock gene expression; FKA protection against H2O2 greater than FKB
43	Puppala et al., 2017	Dihydromethysticin	Mouse lung tumorigenesis (A/J)	In vivo CO2 dosing, DNA isolation, LC-MS/MS quantification, potassium permanganate solution staining, analytical thin layer chromatography, compound visualisation via UV light, mass spectrometry	- Methylenedioxy and lactone functional groups on the chemopreventive activity of DHM used three rationally designed synthetic analogs that can respectively block methylene hydroxylation, lactone hydrolysis, or both routes of metabolism; methylenedioxy functional group of DHM is

					critical for its chemopreventive activity
					while the lactone functional group tolerates
					modifications
					- Compounds 13 and 15, devoid of the
	-				dioxy functional group of DHM while
					retaining the five-membered ring, did not
		\mathbf{O}			show any significant inhibitory activity
		0			against NNK-induced O6 -mG formation or
		07			lung tumor multiplicity in A/J mice; 14,
					with the intact methylenedioxy functional
					group but the mask of the lactone functional
					group, recapitulated the O6 -mG adduct
		σ			reduction potential and antitumorigenic
					efficiency of DHM. The reduction in O6-mG
					correlates well with the blockage of lung
					adenoma formation
					- DHK, which lacks the methylenedioxy
		0			group, is inactive against NNK-induced
		0			DNA adducts
	_				and tumorigenesis
44	Roman	5,6-dehydrokavain	Human glioma (U251),	In vitro	- 5,6-DHK demonstrates highly selective
	Junior et	(Desmethoxyyangoni	human breast	Antiproliferative assays	antiproliferative effect
	al., 2017	n),	adenocarcinoma (MCF-7),		
		Dihydro-5,6-	human ovarian cell		
		dehydrokavain	expressing drugs resistance		

		IUSCript	(NCI-ADR/RES), human renal cell adenocarcinoma (786-o), human large cell lung cancer (NCI-H460), human prostate adenocarcinoma (PC-3), human ovary adenocarcinoma (OVCAR- 03), human colon adenocarcinoma (HT-29), human chronic myeloid		
		R R	leukemia (K-562), immortalized human		
			keratinocytes (HaCat)		
45	Rossette et	Flavokavain B	Human umbilical vein	In vitro	- Antimetastatic action of FKB in vivo might
	al., 2017		endothelial cells (HUVECs),	Cell viability MTT assay	be related to inhibition of angiogenesis.
		Ο	human brain endothelial	Tube formation assay	- Significant inhibition of cell migration at
		Č	cells.	Wound healing assay	low FKB concentration.
	-				- Dose-dependently reduced endothelial cell
			Zebrafish model	In vivo	differentiation into capillary structures
				Zebrafish strain and drug treatment	- Dose dependently inhibited HUVEC tube
					formation
					- Zebrafish model demonstrated dose-
					dependent reduction of subintestinal veins

)t			and intersegmental vessel formation as well as disruption of vascular morphology of SIVs.
46	Seo & Oh, 2013	Non-small Cell Lung Cancer (H1975)	In vitro Proliferation assay, Western Blot	 Dose and time dependently Inhibited cell proliferation Robust degradation of Hsp90's client proteins including EGFR, Met, Her2, Akt, and Cdk4 in a concentration-dependent manner
47	Shaik et Methysticin, al., 2009 Kavain, Dihydrokavain, Dihydromethysticin, Desmethoxyyangonin	Human lung adenocarcinoma (A549)	In vitro Luciferase-based assay, silica gel chromatography, HPLC, H NMR, C NMR, optical rotations, mass spectrometry analysis, western blot analysis	- Kavain, dihydrokavain, DHM, and desmethoxyyangonin are about 100–300 times less active than methysticin
48	Song et al., 2017 Yangonindimer A, B and C	Human lung cancer cell line (NCI-H46), human colorectal adenocarcinoma (SW480), human hepatocellular carcinoma (HepG2)	In vitro MTT assay	- None of the compounds showed significant cellular proliferation inhibition
49	Tang et al., Flavokavain B 2015	Human normal renal cells (HK-2), human acute	In vitro Cell viability assay, Apoptosis assay	 Inhibited proliferation in vivo and in vitro Dose dependently induced intrinsic

			lymphoblastic leukemia	Western blot assay	apoptotic pathway through cleavage and
			(CEM-C1), human acute		activation of caspase 3, and cleavage and
			lymphoblastic leukemia	In vivo	degradation of PARP
		\mathbf{O}	(RS4-11),		- Induced transcription dependent pathway
	-		Balb/c xenograft mice		of apoptosis in vitro and in vivo through
			(CCRF-CEM),	Ex vivo	increasing p53 expression, resulting in the
		\mathbf{O}	human acute lymphoblastic	Isolation of bone marrow specimens from patients	upregulation of pro apoptotic genes Puma
		0	leukemia (B-ALL or T-ALL)		and Bax in vivo and in vitro
		0)			- Dose dependant inhibition of proliferation
					observed in B-ALL and T-ALL but no effect
					on normal cells ex vivo
50		6			
50	Tang et al.,	Flavokavain A,	In vitro	In vitro	- FKB inhibits growth of AR-negative,
	2010	Flavokavain B	Human prostate cancer	MTT assay, Fluorescence-activated cell sorting	hormone refractory PCa cell lines (DU145,
			(LNCaP, LAPC-4, DU145	analysis of apoptosis, Quantification of apoptosis	PC3) ~ 90%, partially reduces growth of
		-	and PC-3), Normal prostate	by ELISA, RT-PCR, RNA interference	AR-positive, hormone sensitive PCa cell
			epithelial (PrECs) and		lines (LNCaP, LAPC4) ~ 32-50%; FKB
		\mathbf{O}	stromal cells (PrSCs)	In vivo	most potent of flavokavains
				Western blot, Immunoprecipitation	- Minimal effects on normal primary prostate
	-		NCR-nu/nu (nude) mice		epithelial and stromal cells
			DU145 xenograft model		- FKB induced apoptosis; activation of
					caspase-3,-8 and -9 activities in AR-negative
					PCa, causes PARP cleavage (DU145 and
					PC-3), increases protein and mRNA
		7			expression of death receptor 5 (DR5) and

		Script			enhances TRAIL ligand induced apoptosis, activates Bax and mitochondrial-mediated apoptotic pathway by upregulation of Bim and Puma and downregulation of XIAP and survivin - FKB inhibits tumor growth in vivo in a DU145 xenograft model and induces Bim expression in tumor tissues
51	Tang et al., 2008	Flavokavain A	Human bladder cancer (T24, UMUC3, TCCSUP, 5637, HT1376, HT1197)	In vitro MTT assay, fluorescence-activated cell sorting analysis of cell cycle distribution, western blot analysis, P27 and p21 Degradation assay, real-time reverse transcription - PCR Immunoprecipitation and kinase assay, SiRNA suppression of p53	 Induced G2-M arrest via suppression of p53 expression by small interfering siRNA in RT4 cells restored Cdc25C expression and down-regulation of p21/WAF1 expression, which allowed Cdc25C and CDK1 activation, which then led to a G2-M arrest and an enhanced growth-inhibitory effect by FKA Selectivity of flavokavain A for inducing a G2-M arrest in p53-defective cells; FKA caused a pronounced CDK1 activation and G2-M arresting p53 knockout but not in p53 wild-type HCT116 cells
52	Yeap et al., 2017	Flavokavain B	Human Cervical cancer (HeLa)	In vitro MTT assay, Flow Cytometry, Gene Expression Profiling Using Microarray, qRT-PCR, Proteome Profiler Antibody Human Cell Stress Array,	 Cytotoxicity (HeLa) Induced G2/M phase arrest Induced apoptosis; loss of membrane potential, differentially regulated mRNA

		anuscript		Superoxide Dismutase (SOD) and Glutathione (GSH) Quantification	expression of cell cycle, apoptosis, cell stress, and MAPK pathways - Upregulation of MAPK pathway, GPx3, HMOX1, CAT from the oxidative pathway, and DDIT3 from the apoptosis pathway, cytochrome C, phospho-p38 alpha (T180/Y182), SOD2, phospho-HSP27 (S78/S82), and HSP70 change - Protects HeLa cells From H2O2-induced cell death, activation of antioxidant neutralizes H2O2-induced ROS, enhanced GSH and SOD Levels
53	Zhao et al., 2011	Flavokavain B	Human oral adenoid cystic carcinoma (ACC-2)	In vitro MTT assay Cell morphology observation and DAPI staining Fluorescence-activated cell sorting analysis of cell cycle distribution Measurement of cytochrome c release from mitochondria RNA interference Western blot analysis RNA isolation and quantitative real-time RT-PCR	 FKB could significantly inhibit the cell proliferation of oral adenoid cystic carcinoma ACC-2 cells; Induced cell-cycle G2-M arrests in ACC-2 cells and subsequent apoptosis in the mitochondrial apoptotic pathway Induced mRNA expression of Bim, Bak, bax, Bcl-2; Bim acted as a potential transcriptional target for FKB
54	Zi &	Flavokavain A, B,	Human bladder cancer (T24,	In vitro	- Kava extract and FKA, FKB, and FKC

RT4, EJ),	Cell viability 3-(4,5-dimethylthiazol-2-yl)-2,5-	cause strong antiproliferative and apoptotic
NCR-nu/nu (nude) mice	diphenyltetrazolium bromide assay, flow	effects in human bladder cancer cells; FKA
	cytometry assays, measurement of cytochrome c	results in cleavage of caspase-3/9 and poly
	release from mitochondria, western blotting and	(ADP-ribose) polymerase in T24 cells in a
	immunoprecipitation, soft agar colony formation	dose- and time-dependent manner. The loss
		of mitochondrial membrane potential and
	In vivo	release of cytochrome c caused by FKA are
		associated with an increase in Bax/Bcl-2
		ratio and Bax confirmation change in T24
		cells
		- FKA decreases the levels of X-linked
		inhibitor of apoptosis and surviving T24
		cells
		- FKA inhibits anchorage-independent
		growth of EJ cells in soft agar and tumour
		growth in nude mice
	RT4, EJ), NCR-nu/nu (nude) mice	RT4, EJ), Cell viability 3-(4,5-dimethylthiazol-2-yl)-2,5- NCR-nu/nu (nude) mice diphenyltetrazolium bromide assay, flow cytometry assays, measurement of cytochrome c release from mitochondria, western blotting and immunoprecipitation, soft agar colony formation In vivo





