



**UNIVERSITI PUTRA MALAYSIA**

**INDUCTION OF APOPTOSIS BY 2',3'-EPOXYISOCAPNOLACTONE  
AND 8-HYDROXYISOCAPNOLACTONE-2',3'- DIOL ISOLATED FROM  
MICROMELUM MINUTUM IN HUMAN T-LYMPHOCYTE LEUKEMIA  
CEM-SS CELLS**

**TAN BOON KEAT.**

**IB 2006 5**

**INDUCTION OF APOPTOSIS BY 2',3'-EPOXY  
ISOCAPNOLACTONE AND 8-HYDROXYISOCAPNOLACTONE-  
2',3'-DIOL ISOLATED FROM *MICROMELUM MINUTUM* IN  
HUMAN T-LYMPHOCYTE LEUKEMIA CEM-SS CELLS**

**TAN BOON KEAT**

**MASTER OF SCIENCE  
UNIVERSITI PUTRA MALAYSIA**

**2006**



**INDUCTION OF APOPTOSIS BY 2',3'-EPOXY  
ISOCAPNOLACTONE AND 8-HYDROXYISOCAPNOLACTONE-  
2',3'-DIOL ISOLATED FROM *MICROMELUM MINUTUM* IN  
HUMAN T-LYMPHOCYTE LEUKEMIA CEM-SS CELLS**

**By**

**TAN BOON KEAT**

**Thesis Submitted to the School of Graduate Studies, Universiti Putra  
Malaysia, in Fulfilment of the Requirement for the  
Degree of Master of Science**

**March 2006**



Abstract of thesis presented to the Senate of Universiti Putra Malaysia in fulfilment of the requirement for the degree of Master of Science

**INDUCTION OF APOPTOSIS BY 2',3'-  
EPOXYISOCAPNOLACTONE AND 8-  
HYDROXYISOCAPNOLACTONE-2',3'-DIOL ISOLATED FROM  
*MIRCAMELUM MINUTUM* IN HUMAN T-LYMPHOCYTE  
LEUKEMIA CEM-SS CELLS**

By

**TAN BOON KEAT**

**March 2006**

**Chairman : Professor Abdul Manaf Ali, PhD**

**Faculty : Institute of Bioscience**

2',3'-epoxyisocapnolactone and 8-hydroxyisocapnolactone-2',3'-diol are two bioactive compounds isolated from the leaves of *Mircamelum minutum*. The cytotoxic effect of the compounds was tested on a variety of human cell lines respectively using MTT assay. They were found to be most sensitive against human T-lymphoblastic leukemia cells (CEM-SS). The inhibition effect of 2',3'-epoxyisocapnolactone and 8-hydroxyisocapnolactone-2',3'-diol at 50% of cell population ( $IC_{50}$ ) was found to be 4.6  $\mu\text{g/ml}$  (13.5  $\mu\text{M}$ ) and 3  $\mu\text{g/ml}$  (7.8  $\mu\text{M}$ ) on CEM-SS cells, respectively. Besides that, the inhibitor effect of the compounds on other human cells were found to be 13.4  $\mu\text{g/ml}$  (39.2  $\mu\text{M}$ ) and 9.0  $\mu\text{g/ml}$  (23.9  $\mu\text{M}$ ) on cervical carcinoma cells (HeLa), 14.2  $\mu\text{g/ml}$  (41.5  $\mu\text{M}$ ) and 7.7  $\mu\text{g/ml}$  (20.5  $\mu\text{M}$ ) on colon adenocarcinoma cells (HT29), 7.4  $\mu\text{g/ml}$  (21.6  $\mu\text{M}$ ) and 5.9  $\mu\text{g/ml}$  (15.7  $\mu\text{M}$ ) on hepatocarcinoma cells (HepG2), 6.5  $\mu\text{g/ml}$  (19.0  $\mu\text{M}$ ) and 7.1  $\mu\text{g/ml}$  (18.9  $\mu\text{M}$ ) on transform liver cells (Chang). For comparative purposes, the  $IC_{50}$



of several clinical cytotoxic drugs against CEM-SS cells were determined. The inhibitor effect of the compounds were more significant compared with methotrexate [ $IC_{50} = >30 \mu\text{g/ml}$  ( $66.1 \mu\text{M}$ )], cytosine arabinoside [ $IC_{50} = >30 \mu\text{g/ml}$  ( $123.5 \mu\text{M}$ )] and colchicines [ $IC_{50} = 8 \mu\text{g/ml}$  ( $20.1 \mu\text{M}$ )]. The compounds also shown near similar  $IC_{50}$  concentration as compare with cis-diamine dichloroplatinum [ $IC_{50} = 3 \mu\text{g/ml}$  ( $10.1 \mu\text{M}$ )], vinorelbine [ $IC_{50} = 3 \mu\text{g/ml}$  ( $3.9 \mu\text{M}$ )] and doxorubicin [ $IC_{50} = 2.4 \mu\text{g/ml}$  ( $4.1 \mu\text{M}$ )]. Furthermore, from proliferation assay study, the compounds were significantly inhibiting the proliferation of cells at  $IC_{50}$  value. From the morphological observation and agarose gel electrophoresis, apoptosis of the compounds on CEM-SS cells was determined. By using phase contrast, fluorescence and electron microcopies, observation on morphological alterations indicating apoptosis was evaluated. From DNA fragmentation, Acridine orange and Propidium iodide staining and DNA content analyses, the compounds were confirmed to have ability in promoting apoptosis. However, the percentage of apoptosis induced is low and the event is time-dependent. At high concentration of  $10 \mu\text{g/m}$ , 2',3'-epoxyisocapnolactone and 8-hydroxyisocapnolactone-2',3'-diol induced necrosis. Furthermore, 8-hydroxyisocapnolactone-2',3'-diol also exhibited better cytotoxicity compared to 2',3'-epoxyisocapnolactone. The induction time for apoptosis by 8-hydroxyisocapnolactone-2',3'-diol in CEM-SS is earlier than 2',3'-epoxyisocapnolactone, which is 4 hours and 12 hours after treatment. Based on the results obtained, 2',3'-epoxyisocapnolactone and 8-hydroxyisocapnolactone-2',3'-diol are able to induced apoptosis.



Abstrak tesis yang dikemukakan kepada Senat Universiti Putra Malaysia sebagai memenuhi keperluan untuk ijazah Master Sains

**KESAN INDUKSI APOPTOSIS OLEH 2',3'-  
EPOXYISOCAPNOLACTONE DAN 8-  
HYDROXYISOCAPNOLACTONE-2',3'-DIOL YANG DIASINGKAN  
DARI *MICROMELUM MINUTUM* KE ATAS JUJUKAN SEL CEM-SS  
T-LYMPHOCYTE LEUKEMIK MANUSIA**

Oleh

**TAN BOON KEAT**

**Mac 2006**

**Pengerusi : Profesor Abdul Manaf Ali, PhD**

**Fakulti : Institusi Biosains**

2',3'-epoxyisocapnolactone dan 8-hydroxyisocapnolactone-2',3'-diol merupakan dua jenis sebatian yang diasingkan dari daun *Micromelum minutum*. Kesan sitotoksik oleh kedua-dua sebatian ke atas pertumbuhan perbagai jenis jujukan kanser sel manusia telah diuji dengan teknik MTT. Mereka didapati lebih sensitive ke atas jujukan sel T-lymphoblastik leukemik manusia (CEM-SS). Kesan perencatan oleh 2',3'-epoxyisocapnolactone and 8-hydroxyisocapnolactone-2',3'-diol pada 50% dari sel populasi (IC<sub>50</sub>) didapati sebanyak 4.6 µg/ml (13.5 µM) dan 3 µg/ml (7.8 µM) masing-masing ke atas sel CEM-SS. Selain itu, kesan perencatan oleh kedua-dua sebatian aktif di atas jujukan sel manusia yang lain juga didapati sebanyak 13.4 µg/ml (39.2 µM) dan 9.0 µg/ml (23.9 µM) ke atas sel servikal karsinoma (HeLa), 14.2 µg/ml (41.5 µM) dan 7.7 µg/ml (20.5 µM) ke atas sel adenokarsinoma usus (HT29), 7.4 µg/ml (21.6 µM) dan 5.9 µg/ml (15.7 µM) ke atas sel hepatokarsinoma (HepG2), 6.5 µg/ml (19.0 µM) dan 7.1



$\mu\text{g/ml}$  ( $18.9 \mu\text{M}$ ) ke atas sel hati yang normal (Chang). Sebagai bandingan,  $\text{IC}_{50}$  dari beberapa jenis ubat pasaran ke atas sel CEM-SS juga diujikan. Kesan perencatan oleh kedua-dua sebatian semulajadi adalah lebih berkesan berbanding dengan ubat-ubatan pasaran seperti methotrexate [ $\text{IC}_{50} = >30 \mu\text{g/ml}$  ( $66.1 \mu\text{M}$ )], cytosine arabinoside [ $\text{IC}_{50} = >30 \mu\text{g/ml}$  ( $123.5 \mu\text{M}$ )] dan colchicines [ $\text{IC}_{50} = 8 \mu\text{g/ml}$  ( $20.1 \mu\text{M}$ )]. Kedua-dua sebatian juga menunjukkan aktiviti  $\text{IC}_{50}$  yang agak sama berbanding dengan cis-diamine dichloroplatinum [ $\text{IC}_{50} = 3 \mu\text{g/ml}$  ( $10.1 \mu\text{M}$ )], vinorelbine [ $\text{IC}_{50} = 3 \mu\text{g/ml}$  ( $3.9 \mu\text{M}$ )] dan doxorubicin [ $\text{IC}_{50} = 2.4 \mu\text{g/ml}$  ( $4.1 \mu\text{M}$ )]. Selain itu, dari pengajian atas teknik pertumbuhan, kedua-dua sebatian adalah ketara dalam perencatan pertumbuhan sel pada  $\text{IC}_{50}$ . Kesan apoptosis bagi kedua-dua sebatian aktif ke atas sel CEM-SS telah diperolehi daripada pemerhatian morfologi dan elektroforesis gel agaros. Dengan menggunakan teknik mikroskop fasa perbezaan, fluorescent dan elektron, pemerhatian ke atas perbezaan morfologi yang berkaitan dengan apoptosis telah dijalankan. Dari keputusan DNA fragmentasi, pewarnaan AO/PI dan pengajian kandungan DNA, kedua-dua sebatian aktif telah menunjukkan kebolehannya dalam menrangsangkan induksi apoptosis. Akan tetapi, peratus bagi induksi apoptosis adalah rendah and kemunculan kesan apoptosis adalah bergantung pada masa rawatan. Bila 2',3'-epoxyisocapnolactone and 8-hydroxyisocapnolactone-2',3'-diol dirawat dengan dos yang tinggi ( $10 \mu\text{g/ml}$ ), ia akan menrangsangkan nekrosis. Selebih daripada itu, 8-hydroxyisocapnolactone-2',3'-diol juga menunjukkan aktiviti sitotoksik yang lebih baik berbanding dengan 2',3'-epoxyisocapnolactone. Kemunculan kesan apoptosis di dalam 8-hydroxyisocapnolactone-2',3'-diol adalah lebih awal berbanding dengan 2',3'-epoxyisocapnolactone, iaitu 4 jam and 12 jam selepas rawatan. Berdasarkan keputusan yang diperolehi, kedua-dua sebatian

semulajadi (2',3'-epoxyisocapnolactone and 8-hydroxyisocapnolactone-2',3'-diol) boleh disimpulkan sebagai agen sitotoksik potensi yang boleh merangsangkan apoptosis.



## ACKNOWLEDGEMENTS

First of all, I would like to express my most sincere thanks and appreciation to my supervisor, Professor Dr. Abdul Manaf bin Ali for his valuable guidance, suggestions, encouragement and advice throughout my project and preparation of the thesis.

I would also like to extend my gratefulness to my co-supervisors; Professor Dr. Mawardi Rahmani and Dr. Noorjahan Alitheen for their support and valuable discussion in making this research a success. Thank you for your kindness to help when I need it most.

Not to be forgotten to express my sincere thanks to Associate Professor Dr. Raha Rahim and Associate Professor Dr. Salmaan Inayat Hussain for their generosity to help and to allow me to use the equipment in their laboratories. Special thanks to Dr. Anthony Ho, his encouragement and patient enlightend me during my difficulties.

I am also very grateful to all my friends in UPM especially to Yih Yih, Ching Ling, Maddie, Yang Ping, Chyan Leong, Dr. Lim, Dr. Khor, Dr Yiap, Varma, Dr. Shuhaimi, Dr. Siti and Dr. Majid. Thank you for willing to share with me your knowledge. Learning from you all is one of the most beautiful things in my life.

Last but not least, my deepest appreciation to my parents who have made me who I am today and my sister, Boon Li and Boon Hooi who have always been there for me.



I certify that an Examination Committee has met on 20<sup>th</sup> March 2006 to conduct the final examination of Tan Boon Keat on his Master of Science thesis entitled “Induction of Apoptosis by 2’,3’-Epoxyisocapnolactone and 8-Hydroxyisocapnolactone-2’,3’-diol Isolated from *Micromelum minutum* in Human T-Lymphocyte Leukemia CEM-SS Cells” in accordance with Universiti Pertanian Malaysia (Higher Degree) Act 1980 and Universiti Pertanian Malaysia (Higher Degree) Regulations 1981. The Committee recommends that the candidate be awarded the relevant degree. Members of the Examination Committee are as follows:

**Fauziah Othman, PhD**

Associate Professor,  
Faculty of Medicine and Health Sciences  
Universiti Putra Malaysia  
(Chairman)

**Johnson Stanslas, PhD**

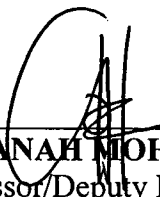
Lecturer,  
Faculty of Medicine and Health Sciences  
Universiti Putra Malaysia  
(Internal Examiner)

**Muhajir Hamid, PhD**

Lecturer,  
Faculty of Biotechnology and Biomolecular Sciences  
Universiti Putra Malaysia  
(Internal Examiner)

**Shaida Fariza Sulaiman, PhD**

Associate Professor  
School of Biological Sciences  
Universiti Sains Malaysia  
(External Examiner)

  
HASANAH MOHD. GHAZALI, PhD  
Professor/Deputy Dean  
School of Graduate Studies  
Universiti Putra Malaysia

Date:

26 APR 2006

This thesis submitted to the Senate of Universiti Putra Malaysia and has been accepted as fulfilment of the requirements for the degree of Master of Science. The members of the Supervisory Committee are as follows:

**ABDUL MANAF ALI, PhD**

Professor  
Faculty of Biotechnology and Biomolecular Sciences  
Universiti Putra Malaysia  
(Chairman)

**MAWARDI RAHMANI, PhD**

Professor  
Faculty of Sciences  
Universiti Putra Malaysia  
(Member)

**NOORJAHAN ALITHEEN, PhD**

Lecturer  
Faculty of Biotechnology and Biomolecular Sciences  
Universiti Putra Malaysia  
(Member)



---

**AINI IDERIS, PhD**  
Professor/Dean  
School of Graduate Studies  
Universiti Putra Malaysia

Date: **11 MAY 2006**



## DECLARATION

I hereby declare that the thesis is based on my original work except for quotations and citations, which have been duly acknowledged. I also declare that it has not been previously or concurrently submitted for any other degree at UPM or other institutions.



**TAN BOON KEAT**

Date: 24 / 4 / 06

## TABLE OF CONTENTS

<b>ABSTRACT</b>	ii
<b>ABSTRAK</b>	iv
<b>ACKNOWLEDGEMENTS</b>	vii
<b>APPROVAL</b>	viii
<b>DECLARATION</b>	x
<b>LIST OF TABLES</b>	xiii
<b>LIST OF FIGURES</b>	xiv
<b>LIST OF ABBREVIATION</b>	xvii

### CHAPTER

1	INTRODUCTION	1
2	LITERATURE REVIEW	5
2.1	Cancer	5
2.1.1	Biology of Tumor	6
2.1.2	Classification of Cancer	6
2.1.3	Leukemia	7
2.2	Chemotherapy	9
2.2.1	Herbal Medicine	10
2.2.2	<i>Micromelum minutum</i>	11
2.3	Terminology of Cell Death	13
2.4	Necrosis	13
2.5	Apoptosis	14
2.5.1	Morphology of Apoptosis	16
2.5.2	Biochemical Regulation of Apoptosis	20
2.5.2.1	Endonuclease Activation	20
2.5.2.2	Phagocytosis	23
2.5.2.3	Intracellular Signaling	24
2.5.2.4	Caspase	27
2.5.3	Molecular Regulation of Apoptosis	33
2.5.3.1	Bcl-2 Family	33
2.5.3.2	Tumor Suppressor Genes	36
3	MATERIALS AND METHODS	40
3.1	Natural compounds	40
3.2	Cell lines	40
3.3	Cells reviving	41
3.4	Cell Line Maintenance	41
3.5	Cytotoxicity Assay	42
3.5.1	MTT Assay	44
3.6	Proliferation Assay	45
3.7	Morphology Assessment	45
3.7.1	Phase Contrast Microscopy	45
3.7.2	Fluorescent Microscopy	46



	[Acridine Orange/ Propidium Iodide (AO/PI) Staining]	
	3.7.3 Electron Microscopy	47
	[Scanning Electron Microscopy (SEM) – (JEOL-JSM 6400)]	
3.8	DNA Fragmentation Assay	48
3.9	Flow Cytometric Analyses	49
3.10	Statistical Analyses	50
4	RESULTS AND DISCUSSION	51
4.1	MTT Assay	51
4.2	Proliferation Assay	61
4.3	DNA Fragmentation Assay	67
4.4	Morphological Assessment of Cell Death	69
	4.4.1 Phase Contrast Microscopy	69
	4.4.2 Fluorescence Microscopy	74
	4.4.3 Electron Microscopy	84
4.5	Cell Cycle Analyses	88
5	CONCLUSION	94
	REFERENCES	97
	BIODATA OF THE AUTHOR	107



## LIST OF TABLES

Table	Page
1 Differential features and significance of necrosis and apoptosis	19
2 The Caspase Family	32
3 Serial dilution gradient	43
4 The inhibition concentration of 50% (IC <sub>50</sub> ) of both natural compounds isolated from <i>Mircomelum minutum</i> (2',3'-epoxyisocapnolactone and 8-hydroxyisocapnolactone-2',3'-diol) against various cell lines determined by using MTT assay	59
5 The inhibition concentration of 50% (IC <sub>50</sub> ) of various standard cytotoxic compounds against CEM-SS cells determined by using MTT assay	59
6 The percentage of CEM-SS cells distribution in different cell cycle phase for untreated cells, cells treated with 2',3'-epoxyisocapnolactone in IC <sub>50</sub> and cells treated with 8-hydroxyisocapnolactone-2',3'-diol in IC <sub>50</sub> by using flow cytometry analysis	93



## LIST OF FIGURES

Figure	Page
1 The chemical structure of 2',3'-epoxyisocapnolactone isolated from the leaves of <i>Micromelum minutum</i>	12
2 The chemical structure of 8-hydroxyisocapnolactone-2',3'-diol isolated from the leaves of <i>Micromelum minutum</i>	12
3 Illustration of the morphological features of necrosis and apoptosis	18
4 Distinct pathways to apoptosis converge on activation of caspase	31
5 The intrinsic and extrinsic pathways leading to apoptosis	35
6 Molecular structure of MTT and formazan	53
7 Cytotoxic activity of 2',3'-epoxyisocapnolactone and 8-hydroxyisocapnolactone-2',3'-diol against CEM-SS cells after 72 hours of incubation. Viability was determined with MTT Assay and data represents mean of triplicate $\pm$ SD	54
8 Cytotoxic activity of 2',3'-epoxyisocapnolactone and 8-hydroxyisocapnolactone-2',3'-diol against HT-29 cells after 72 hours of incubation. Viability was determined with MTT Assay and data represents mean of triplicate $\pm$ SD	55
9 Cytotoxic activity of 2',3'-epoxyisocapnolactone and 8-hydroxyisocapnolactone-2',3'-diol against HeLa cells after 72 hours of incubation. Viability was determined with MTT Assay and data represents mean of triplicate $\pm$ SD	56
10 Cytotoxic activity of 2',3'-epoxyisocapnolactone and 8-hydroxyisocapnolactone-2',3'-diol against HepG2 cells after 72 hours of incubation. Viability was determined with MTT Assay and data represents mean of triplicate $\pm$ SD	57
11 Cytotoxic activity of 2',3'-epoxyisocapnolactone and 8-hydroxyisocapnolactone-2',3'-diol against Chang cells after 72 hours of incubation. Viability was determined with MTT Assay and data represents mean of triplicate $\pm$ SD	58
12 The cytotoxic effect of 2',3'-epoxyisocapnolactone on CEM-SS cell concentration compared with untreated cell. Data represents mean of triplicate $\pm$ SD	65





<b>Figure</b>	<b>Page</b>
13 The cytotoxic effect of 8-hydroxyisocapnolactone-2',3'-diol on CEM-SS cell concentration compared with untreated cell. Data represents mean of triplicate $\pm$ SD	66
14 Effect of 2',3'-epoxyisocapnolactone and 8-hydroxyisocapnolactone-2',3'-diol on DNA fragmentation	68
15 Phase contrast microscopy examination of untreated CEM-SS cells	71
16 Phase contrast microscopy examination of CEM-SS cells treated with 2',3'-epoxyisocapnolactone	72
17 Phase contrast microscopy examination of CEM-SS cells treated with 8-hydroxyisocapnolactone-2',3'-diol	73
18 Fluorescence microscopy examination of untreated CEM-SS cells	78
19 Fluorescence microscopy examination of CEM-SS cells treated with 2',3'-epoxyisocapnolactone	79
20 Fluorescence microscopy examination of CEM-SS cells treated with 8-hydroxyisocapnolactone-2',3'-diol	80
21 The percentage of viable, apoptotic and necrotic CEM-SS cell in the population after treated with 2',3'-epoxyisocapnolactone at various time courses	81
22 The percentage of viable, apoptotic and necrotic CEM-SS cell in the population after treated with 8-hydroxyisocapnolactone-2',3'-diol at various time courses	82
23 The percentage of viable, apoptotic and necrotic CEM-SS cell in the untreated population and population after treated with 0.1% DMSO at various time courses	83
24 Scanning electron microscopy examination of untreated CEM-SS cells	85
25 Scanning electron microscopy examination of CEM-SS cells treated with 2',3'-epoxyisocapnolactone	86
26 Scanning electron microscopy examination of CEM-SS cells treated with 8-hydroxyisocapnolactone-2',3'-diol	87



<b>Figure</b>		<b>Page</b>
27	DNA fluorescence histograms of the effect of 2',3'-epoxyisocapnolactone on the progression through the cell cycle	91
28	DNA fluorescence histograms of the effect of 8-hydroxyisocapnolactone-2',3'-diol on the progression through the cell cycle	92



## LIST OF ABBREVIATIONS

%	Percentage
ALL	Acute Lymphocytic Leukemia
ANLL	Acute Non-Lymphocytic Leukemia
AO	Acridine Orange
ATCC	American Type Culture Collection
ATP	Adenosine triphosphate
bp	Base pair
CEM-SS	T-Lymphoblastic Leukemia
CGM	Complete Growth Medium
Chang	Normal Liver Cell
CO <sub>2</sub>	Carbon dioxide
Da	Dalton
DMSO	Dimethylsulfoxide
DNA	Deoxyribonucleic acid
EDTA	Ethylenediaminetetraacetic acid Di-Sodium Salt
ELISA	Enzyme linked immunosorbent assay
FBS	Fetal Bovine Serum
HeLa	Cervical Carcinoma
HepG2	Hepato Carcinoma
HT29	Colon Carcinoma
IC <sub>50</sub>	Inhibition Concentration 50%
mg	Milligram
ml	Milliliter



MTT	3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide
nm	Nanometer
O.D.	Optical Density
PBS	Phosphate Buffered Saline
PI	Propidium Iodide
RNA	Ribonucleic acid
RPM	Rotation per Minute
SDS	Sodium dodecyl sulfate
SEM	Scanning Electron Microscope
UV	Ultra violet
μg	Microgram
μl	Microliter



# CHAPTER 1

## INTRODUCTION

Cancer chemotherapy, the treatment or control of cancer using anticancer drugs which are highly toxic medications that destroy cancer cells by interfering with their growth or preventing their reproduction (Altman and Sarg, 1992). It has played a role in cancer treatment for almost half a century. Years of testing and research have proved chemotherapy to be an effective cancer treatment. It may be the only treatment, or it may be used in combination with other treatments, including surgery and radiation therapy. Chemotherapy works by killing rapidly dividing cells. These cells include cancer cells, which continuously divide to produce more cells, and healthy cells that divide quickly, such as those in bone marrow, gastrointestinal tract, reproductive system and hair follicles. Healthy cells usually recover shortly after chemotherapy is complete (Mayo Foundation for Medical Education and Research, 2003). It differs from surgery or radiation in that it is always used as a systemic treatment. In chemotherapy the medicines travel throughout the whole body or system rather than being confined or localized to one area such as the breast, lung, or colon. Thus chemotherapy can reach cancer cells that may have spread to other parts of the body.

More than 100 drugs are currently used for chemotherapy, either alone or in combination. Many more are expected to become available. These chemotherapy medicines are vary widely in their chemical composition, how they are taken, their usefulness in treating



specific forms of cancer, and their side effects. New medications are first developed through laboratory research in test tubes and animals. Then, their safety and effectiveness are tested for clinical trials in humans (American Cancer Society, 2001).

Chemotherapy drugs are divided into several categories based on how they affect specific chemical substances within cancer cells, which cellular activities or processes the drug interferes with, and which specific phases of the cell cycle the drug affects. Most chemotherapy is given as a combination of drugs that work together to kill cancer cells. Some of the types of chemotherapy medications commonly used to treat cancer include:

- 1) Alkylating agents. These medications interfere with the growth of cancer cells by blocking the replication of DNA. Examples of alkylating agents include busulfan, cisplatin, carboplatin, chlorambucil, cyclophosphamide, ifosfamide, dacarbazine (DTIC), mechlorethamine (nitrogen mustard), and melphalan.
- 2) Antimetabolites. These drugs block the enzymes needed by cancer cells to live and grow. Examples of antimetabolites include 5-fluorouracil, capecitabine, methotrexate, gemcitabine, cytarabine (ara-C), and fludarabine.
- 3) Antitumor antibiotics. These antibiotics are different from those used to treat bacterial infections. It interferes with DNA, blocking certain enzymes and cell division, and changing cell membranes. Examples of antitumor antibiotics include dactinomycin, daunorubicin, doxorubicin (Adriamycin), idarubicin, and mitoxantrone.
- 4) Mitotic inhibitors. These drugs inhibit cell division or hinder certain enzymes necessary in the cell reproduction process. Examples of mitotic inhibitors include paclitaxel, docetaxel, etoposide (VP-16), vinblastine, vincristine, and vinorelbine.
- 5) Nitrosoureas.



These medications impede the enzymes that help repair DNA. Examples of nitrosoureas include carmustine (BCNU) and lomustine (CCNU) (American Cancer Society, 2001).

Previous studies have demonstrated that a wide range of anticancer agents, including chemotherapeutic agents, hormones, and various biologicals, induce apoptosis in malignant cells *in vitro* (Mesner *et al*, 1997; Kaufmann and Earnshaw, 2000). It is important to emphasize that this treatment-induced apoptosis is not merely a tissue culture phenomenon. Serial examination of peripheral blood mononuclear cells from acute leukemia patients undergoing induction therapy has demonstrated that various agents, including cytarabine, mitoxantrone, etoposide, paclitaxel, and topotecan, cause a marked increase in the number of apoptotic blasts (Li *et al*, 1994). Characteristic apoptotic changes have also been described in solid tumors after treatment of mice with various cytotoxic drugs, including cytarabine, 5-fluorouracil (5FU), fludarabine, doxorubicin, cyclophosphamide, cisplatin, etoposide, dactinomycin, and camptothecin (Kaufmann and Earnshaw, 2000).

For chemotherapy, natural products have been important sources of medicines for many traditional communities around the world. Natural products or their structural relatives comprise about 50% of the drugs that are used in cancer chemotherapy (Mann, 2002). In Malaysia, out of 12,000 species of higher plants which are found in this country, that are more than 1,000 species are said to have therapeutic properties and currently being used in the local traditional medicine system (Said, 1995). According to another report by Burkill in 1966, there are about 6,000-7,000 species of higher plants that have been



reported to have therapeutic or medicinal properties in Peninsular Malaysia and its surrounding islands (Burkill, 1966). Since the use of natural products in cancer chemotherapy has grown tremendously, the study of mechanism and mode of action of plant extracts become more and more important. In many research, plant related derivatives can be synthesized by knowing their biochemical reaction against cancer, and these derivatives can even have greater effect over the original compounds.

In this study, two natural compounds, 2',3'-epoxyisocapnolactone and 8-hydroxyisocapnolactone-2',3'-diol, which were isolated from the leaves of *Micromelum minutum*, were tested for their cytotoxic activity against human T-lymphoblastic leukemia cells (CEMSS)

The objectives of this study were:

- To evaluate the cytotoxic and antiproliferative activities of 2',3'-epoxyisocapnolactone and 8-hydroxyisocapnolactone-2',3'-diol against human T-lymphoblastic leukemic cells (CEM-SS) in terms of proliferation, morphological changes and the mode of cell death.
- To investigate the induction of apoptosis by 2',3'-epoxyisocapnolactone and 8-hydroxyisocapnolactone-2',3'-diol in treated CEM-SS cells.



## CHAPTER 2

### LITERATURE REVIEW

#### 2.1 Cancer

Cancer, a general term for more than 100 diseases which characterized by an uncontrolled, abnormal growth of cells appear in different parts of the body that can spread to other parts of the body (Altman and Sarg, 1992). It is a potentially fatal disease caused mainly by environmental factors that mutate genes encoding critical cell-regulation proteins. The resultant aberrant cell behavior leads to expensive masses of abnormal cells that destroy surrounding normal tissue and can spread to vital organs resulting in disseminated, commonly a harbinger of imminent patient death (Alison, 2002). Cancer cells contain many genetic alterations that accumulate as tumors develop. Over the last 20 years, considerable information has been gathered on regulation of cell growth and proliferation leading to the identification of the involvement of specific genes at the molecular level (Macdonald and Ford, 1997).

The incident of cancer is rising with doubling in new cancer cases and cancers related deaths expected over the next 20 years (Alison, 2002). Part of the reason for this rise is that life expectancy is steadily rising and most cancers are more common in an ageing population. More significantly, a globalization of unhealthy lifestyles, particularly