

**Figure 2.** Summary of differentiation of the *rpoB* gene of *M. tuberculosis* complex from other mycobacteria and pathogenic bacteria using a combination of PCR-RFLP pattern of regions E and D.

**Table 4** Summary of PCR amplification of 9 *rpoB* gene fragments in bacteria.

	A	B	C	D	E	F	G	H	I
	36-1073	1024-1666	1615-2611	2335-2492	2335-2962	2856-3645	3216-4028	3935-4357	4261-4652
1. <i>C. diphtheriae</i>	-	-	-	+	-	-	-	+	-
2. <i>E. coli</i>	-	-	-	+	-	-	-	+	-
3. <i>H. influenza</i>	-	-	-	+	-	-	-	+	-
4. <i>K. pneumoniae</i>	-	-	-	+	-	-	-	+	-
5. <i>N. meningitidis</i>	-	-	-	+	-	-	-	+	-
6. <i>Ps. aeruginosa</i>	-	-	-	+	-	-	-	+	-
7. <i>Ps. pseudomallei</i>	-	-	-	+	-	-	-	+	-
8. <i>S. aureus</i>	-	-	-	+	-	-	-	+	-
9. <i>S. epidermidis</i>	-	-	-	+	-	-	-	+	-
10. <i>S. pyogenes</i>	-	-	-	+	-	-	-	+	-
11. <i>S. viridans</i>	-	-	-	+	-	-	-	+	-

+ = amplified; - = not amplified

**Table 5** Summary of PCR amplification of 9 *rpoB* gene fragments in mycobacteria.

	A	B	C	D	E	F	G	H	I
	36-1073	1024-1666	1615-2611	2335-2492	2335-2962	2856-3645	3216-4028	3935-4357	4261-4652
1. <i>M. tuberculosis</i>	+	+	+	+	+	+	+	+	+
2. <i>M. africanum</i>	+	+	+	+	+	+	+	+	+
3. <i>M. bovis</i> BCG	+	+	+	+	+	+	+	+	+
4. <i>M. microti</i>	-	+	+	+	+	+	-	+	+
5. <i>M. avium</i>	-	-	+	+	+	-	+	+	+
6. <i>M. chelonae</i>	-	-	-	+	-	-	-	+	-
7. <i>M. duvalii</i>	-	-	ND	+	ND	-	-	+	ND
8. <i>M. goodii</i>	-	-	-	+	+	-	+	+	+
9. <i>M. kansasii</i>	-	-	-	+	+	-	+	+	-
10. <i>M. phlei</i>	-	-	ND	+	ND	-	-	+	ND
11. <i>M. scrofulaceum</i>	-	-	+	+	+	-	-	+	-
12. <i>M. smegmatis</i>	-	-	ND	+	ND	-	-	+	ND
13. <i>M. vaccae</i>	-	-	ND	+	ND	-	-	+	ND
14. <i>M. intracellulare</i>	-	-	+	+	+	-	+	+	+

**+** = amplified; **-** = not amplified; **ND** = not determined

## เนื้อหางานวิจัย

### CATALASE-GENETICALLY MODIFIED *LACTOBACILLUS PLANTARUM* AS A STARTER CULTURE IN FERMENTED MEAT PRODUCT

#### ABSTRACT

The property of catalase is very rare in lactobacilli but desirable for starter cultures as it prevents food product defects caused by hydrogen peroxide. *Lactobacillus sake* SR911 is a local strain which produces a heme dependent catalase. The catalase gene of *Lactobacillus sake* SR911 was amplified by polymerase chain reaction and cloned in pMEx8 vector and introduced into *E.coli* UM2. The catalase gene was subcloned into *E.coli-lactococcus* shuttle vector pGKV210 and introduced into *Lactobacillus plantarum* a catalase negative strain isolated from Thai fermented sausage(NHAM). The transformed lactobacilli were shown to decompose hydrogen peroxide and the molecular size of the recombinant protein was 65,000 Da on SDS-PAGE. The DNA sequence contains 1637 bp encoding 478 amino acids. The level of expression of catalase gene was increased when expressed under strong lactococcus promoter p59 in vector pIL1020. The transformed strain showed stable catalase activity and the recombinant plasmid was retained in more than 95% of the cell after 20 generations. The stability of recombinant plasmid was also investigated when using as starter culture in the process of meat fermentation. This catalase-genetically modified strain may prove to be useful as a starter culture in fermented meat product.

#### INTRODUCTION

Lactobacilli are important microorganisms and are widely used in food and agricultural industries. Their main function is to produce acid which lowers the pH thus prevents the growth of spoilage bacteria in the fermented product. They are also involved in the development of flavor and texture in the final product (Mckay et.al., 1990). Their significant roles in food conversion and preservation have provided the opportunity to develop new strains with specialized role. In the fermentation process the presence of oxygen may occur and lactic acid bacteria may produce hydrogen peroxides as their common metabolites (Whittenbury et.al., 1964). Hydrogen peroxide is a very strong oxidizing agent and if accumulated may affect the sensory quality of the product as a result of rancidity or color defects (Niven et.al., 1957). Lactobacilli are commonly considered as

lacking of catalase until recently whereby many studies have shown that lactic acid bacteria contain at least 2 groups of catalase (Wolf et.al., 1991; Engesser et.al., 1994). The two groups of catalase include true catalase or heme catalase and the non-heme catalase or pseudocatalase or manganese catalase. The first group of enzyme can be detected in many species of lactic acid bacteria if heme or hematin is present with different activities. In the second group, the enzyme is active without addition of heme and is found in very few species (Engesser et.al., 1994). The *kata* gene encoding the true catalase from *L. sake* LTH677 and *L. sake* SR911 has been cloned and characterized (Knauf et.al., 1992; Noonpakdee et.al., 1996). Recently the pseudocatalase gene from *Lactococcus plantarum* has been also cloned and characterized (Igarashi et. al., 1996).

*Lactobacillus* sp. TISTR891 and *Lactobacillus plantarum* TISTR850 are isolated from local fermented meat products (NHAM) and are catalase negative. *Lactobacillus sake* SR911 on the contrary contains high heme dependent catalase. Since catalase is one of the desired properties for starter cultures used in food fermentation as it can prevent flavor and color defects in the products due to the effect of hydrogen peroxide (Lucke et.al., 1985), the transfer of the catalase gene into these lactobacilli can give them the ability to produce catalase and simplify starter preparations (Hammes et.al., 1994). Moreover the catalase gene may be used as food grade marker since detection of oxygen producing colonies is very simple. In this study, we report the cloning and transferring of the catalase gene from *L. sake* SR911 into selected lactic acid bacteria that lack this enzyme. The work presented in this report demonstrates a successful cloning and expression of the catalase gene into *Lactobacillus* sp. TISTR891 and *L. plantarum* TISTR 850 using an *E.coli*-*lactococcal* shuttle vector pGKV210 and expression vector pIL1020. The amount of catalase activity obtained from these strains were compared. Study on the properties of transformed *L.plantarum* TISTR 850 as starter culture in food was also investigated.

## **MATERIALS AND METHODS**

### **Bacterial Strains and Plasmids**

The bacterial strains and plasmids used in this study are listed in Table 1.

### **Media and Growth Conditions**

The *E. coli* and lactobacilli strains were grown in LB medium (Davies et.al., 1980) and MRS medium (DeMan et.al., 1960) as previously described (Tanasupawat et. al., 1993). To detect catalase activity in lactobacilli, cells were grown in MRS media in the presence of hematin at a final concentration of 30  $\mu$ M as previously described (Wolf et.al.,

1991). Selective media containing ampicillin at 100 µg/ml or erythromycin at 200 µg/ml were used for selection of *E. coli* and 20 µg /ml erythromycin was used for selection of lactobacilli.

### DNA Preparation

Chromosomal DNA from *L. sake* was isolated according to the method previously described (Anderson et.al., 1983). Proteinase K was used at the final concentration of 20 µg/ml before extraction with phenol/chloroform/isoamyl alcohol. Plasmid DNA of *E. coli* was isolated by standard alkaline lysis method (Sambrook et.al., 1989). Plasmid DNA of lactobacilli were isolated according to the method described by Anderson and McKay (Anderson et.al., 1983).

### Enzymes

DNA-modifying enzymes supplied with appropriate buffers were obtained from Boehringer Mannheim. *Hind* III cut λ DNA was used as molecular weight marker.

### Cloning Procedure

The catalase gene including its promoter region was amplified from chromosomal DNA of *L. sake* SR911 by polymerase chain reaction. PCR amplification was carried out in 50 µl mixture containing 5 µl of 10x PCR buffer (10 mM KCl, 10 mM ammonium sulfate, 20 mM Tris-HCl (pH 8.8 at 25 °C), 2 mM Mg<sub>2</sub>SO<sub>4</sub> and 0.1% Tritonx-100), 1 unit of Vent<sup>R</sup> DNA Polymerase (New England Biolabs, Beverly, Massachusetts, U.S.A) and 200 µM each of the deoxynucleoside triphosphate. The amplification was performed on a DNA thermal cycler (Perkins Elmer Cetus Model TCI) using the following programme; denaturation (1 min at 90 °C) ; annealing (1 min at 55° C) and extension (2 min at 72° C) for 29 cycles. The final cycle consisted of the following steps, denaturation (1 min at 90 °C), annealing (1 min at 55° C) and extension (5 min at 72 °C). PCR primers were designed from -35 region of the gene encoding catalase of *L. sake* LTH677 as follow

oligo1, 5'primer 5' CGC GGA TCC GAG CAA GCT CTT CAT TGACG 3'

- 3 5

oligo2, 3'primer 5' AAA ACT GCA GAG ATA ACG AAT AAA AAA TAG GTG 3'

The amplified product was gel purified and ligated into pUC19 which had been digested with *Sma* I. The resulting recombinant plasmid referred as pUCK was subcloned

into the *E.coli-lactococcus* shuttle vector pGKV210 yielding pGKV8E. The amplified product was also ligated to pIL1020 vector which had been digested with *Eco* RV to yield the plasmid pILK with two possible orientations of the catalase gene under the control of p59 lactococcal promoter.

#### **Transformation of *E. coli* UM2 and lactobacilli.**

Competent *E. coli* cells prepared by  $\text{CaCl}_2$  method (Sambrook et.al., 1989) were transformed and transformants were screened by plating on LB medium with 100  $\mu\text{g/ml}$  of ampicillin or 200  $\mu\text{g/ml}$  erythromycin. All antibiotic resistant transformants were further analyzed for expression of the catalase. The presence of plasmids with the catalase gene cassette were analysed by restriction digestion. Transformation of *Lactobacillus* sp. TISTR891 and *L. plantarum* TISTR850 were carried out by electroporation according to the protocol previously described using Biorad Gene Pulser in a 0.2 cm cuvette with the field strength of 1.25 kV/cm (Bringell et.al., 1990; Thompson et.al.,1996).

#### **Preparation of cell-free extracts from lactobacilli**

To determine the catalase activity in crude cell extracts, aliquots of overnight culture (30ml) were harvested by centrifugation at  $4^\circ\text{C}$  and washed with ice-cold phosphate buffer (50mM, pH 7). The washed cells were suspended in the same buffer with 0.5 mM phenylmethylsulfonyl fluoride and sonicated for three 1-min periods in an ice bath. After centrifugation, the crude cell extracts were assayed immediately. The protein concentrations were determined by Bio-Rad protein assay.

#### **Catalase activity test**

Catalase positive clones were detected by flooding colonies on agar plate with hydrogen peroxide solution (0.87 M) and the bubble producing colonies were selected for further analysis. The quantitative assay for the presence of catalase in bacterial cells was performed according to the method previously described (Sinha, 1972; Noonpakdee et.al., 1996). Total cellular catalase activity was expressed as  $\mu\text{mole}$  of hydrogen peroxide decomposed per minute per  $3 \times 10^8$  CFU. The specific activity of catalase in crude cell extract was expressed as  $\mu\text{mole}$  of hydrogen peroxide decomposed per minute per mg protein.

### **Protein pattern analysis**

Total soluble proteins from *E. coli* UM2 and *Lactobacilli* were extracted and resolved by SDS-polyacrylamide gel electrophoresis (Laemmli, 1970). Standard protein markers from Bio-Rad were used as molecular weight markers.

### **In-situ activity staining of the catalase.**

Total soluble proteins were isolated as described above and separated on a 7.5 % nondenaturing polyacrylamide gel. The gel was stained using ferricyanide according to the method previously described (Wayne et.al., 1986).

### **DNA sequencing**

DNA sequencing was achieved by using automated sequencing analyzer (Perkin Elmer ABI No377) with ABI Prism Dye Terminator Cycle Sequencing Ready kit. Reactions were performed as described by the instructions of the manufacturer. Subclones were constructed in pUC19, and the insert was sequenced from both sides by using universal primer and or reverse oligonucleotides primer. DNA sequence and amino acid sequence were analyzed with PCgene software Release Version 6.8.

### **Preparation of fermented meat product(NHAM)**

To prepare fermented meat product(NHAM) , Culture of *L. plantarum* TISTR 850 or *L. plantarum* TISTR 850 harboring the recombinant plasmid was inoculated into mixture of ground pork, minced skin pork, rice and garlic at concentration of  $10^8$  CFU/g , then incubated at 30 C for 36-48 h. As a control, no culture was inoculated.

### **Lipid extraction method**

Lipid extraction of fermented meat product was extracted according to Bligh-Dyer method (Bligh and Dyer,1959).

### **Iodine number value determination**

Iodine number of the fermented meat product was determined according to Wijs method (AOAC:official method,1995)

## **RESULTS**

Cloning of the catalase gene from *L. sake* SR911 in pGKV210 and pIL1020

The catalase gene of *L. sake* SR911 including its promoter region was amplified using the primers derived from the nucleotide sequence of *L. sake* LTH677 (Knauf et.al., 1992). The catalase gene from *L. sake* SR911 was cloned as a 1.6 kb amplified fragment into *Sma* I cut pUC19. The resulting plasmid was designated pUCK 233 and then

introduced into *E. coli* UM2 which is catalase negative as a result of mutation of *KatG* and *KatE*. Three transformants were selected on LB agar plate containing ampicillin and expression of catalase was revealed by flooding the colonies with hydrogen peroxide. All transformants were found to contain a 1.6 kb size of catalase gene in plasmid DNA and exhibited the same level of catalase activity as measured by quantitative catalase assay. One of the recombinant plasmid was subcloned into *E.coli-lactococcus* shuttle vector pGKV210 resulting in plasmid pGKV8E and later introduced into *Lactobacillus sp.* TISTR891 by electroporation. The catalase activity produced by transformed lactobacilli must be due to expression of the catalase gene under the control of its own promoter since *Lactobacillus sp.* TISTR891 is catalase negative. Plasmid pGKV210 is a general cloning vector which contained the origin of replication pWV01. All transformants grown in erythromycin plate were found to be catalase positive. In order to increase the catalase activity, we also cloned this amplified 1.6 kb DNA fragment into *EcoR* V cut pIL1020, an expression vector which contained the same origin of replication of pWV01 and the strong lactococcal promoter p59 (Van der Vossen et.al., 1985). The ligation mixture was used to transform *E. coli* UM2 as intermediate host. More than 300 erythromycin resistant transformants were screened for catalase activity. Three clones were found to be catalase positive. After restriction analysis of the purified plasmid DNA, we found that these three recombinant plasmids contained the same size 1.6 kb insert in either of the two possible orientations. Plasmid from one transformant designated pILK01 had the catalase gene cloned downstream from the p59 promoter. Two recombinant plasmid designated pILK02 and pILK03 had the catalase gene cloned in the opposite direction from p59 promoter (figure1). All 3 recombinant plasmids were then used to transform *Lactobacillus sp.* TISTR891 and pILK01 was also used to transform *Lactobacillus plantarum* TISTR850 by electroporation.

Expression of the catalase gene of *Lactobacillus sake* SR911 from pGKV8E, pILK01, pILK02 and pILK03 in *E. coli* UM2

All transformed *E. coli* UM2 harboring plasmid pGKV8E, pILK01, pILK02, and pILK03 exhibited catalase activities by quantitative assay as compared to control *E. coli* UM2 host strain and *E. coli* UM2 with shuttle vectors pGKV210, pIL1020. The expression of catalase in transformed *E. coli* UM2 did not depend on exogenous hematin unlike in *L. sake* SR911 which we had to supply hematin in the growth medium before catalase assay



suggesting that *E. coli* strain can synthesize porphyrin group (heme) unlike most lactic acid bacteria (Whittenbury et.al., 1964).

The protein expressed from the 1.6 kb amplified DNA insert in *E. coli* UM2 using pGKV8E, pILK01, pILK02 and pILK03 were analysed on denaturing 7.5 % SDS polyacrylamide gel (SDS-PAGE) and the result is shown in Figure 2. In *E. coli* UM2 bearing the recombinant plasmids, the protein patterns showed the presence of an additional protein band with the molecular size of approximately 62,000 Da (lane 3, 5, 6 and 7). This protein band was not present in *E. coli* UM2 with vector pGKV210 or expression vector pIL1020 (lane 2, 4). The protein of the same molecular size was also observed in the protein pattern of wild type catalase strain *L. sake* SR911 (lane 1). This apparent recombinant protein is most likely to be encoded by the gene carried on the plasmids pGKV8E, pILK01, pILK02 and pILK03 since *E. coli* UM2 host is catalase negative. The intensity of protein band from transformed *E. coli* harboring pILK01 (lane 5) is approximately 2 fold (as determined by scanning with a densitometer) as compared with that of transformed *E. coli* cells harboring pGKV8E, pILK02 and pILK03 (lane 3, 6, 7).

Expression of the catalase gene of *L. sake* SR911 from pGKV8E, pILK01, pILK02 and pILK03 in *Lactobacillus sp.* TISTR891 and *Lactobacillus plantarum* TISTR850

The 1.6 kb insert of the catalase gene in the recombinant plasmid pGKV8E, pILK02 and pILK03 were expressed under its own promoter. The pILK01 however, had the catalase gene expressed under the transcriptional control of the lactococcal promoter p59. The expression of these different constructs was evaluated in the transformed lactobacilli by quantitative catalase assay and by in-situ activity staining in polyacrylamide gel electrophoresis. The data in table 2 showed that all transformed lactobacilli with recombinant plasmids exhibited heme dependent catalase activity. However, *Lactobacillus sp.* TISTR891 and *L. plantarum* TISTR850 host strains with and without the plasmids showed no activity at all. The specific catalase activity from transformed *Lactobacillus sp.* TISTR891 harboring pILK01 was approximately 2 times higher than those from *Lactobacillus sp.* TISTR891 harboring pGKV8E, pILK02 and pILK03. Therefore, the expression of the catalase gene under the transcriptional control of lactococcus promoter p59 was markedly increased compared to the expression under the control of its own promoter. Plasmid pILK01 was then used to transform another catalase negative strain, *Lactobacillus plantarum* TISTR850. The specific catalase activity of transformed *L. plantarum* harboring pILK01 increased almost 3 times higher than that of *Lactobacillus sp.*

TISTR891 harbouring plasmid pILK01 and about 6 times higher than that of *Lactobacillus* sp. TISTR891 harbouring pILK02 and pILK03. The catalase activity in transformed *L. plantarum* TISTR850 was about 3 times higher than that of wild type *L. sake* SR911.

The result of catalase in-situ activity staining on a non-denaturing polyacrylamide gel was shown in Figure 3. The pattern of catalase protein bands from transformed lactobacilli with recombinant plasmids (lane 3, 5, 6, 8), was similar to that of wild type *L. sake* SR911 (lane 1). The intensity of the recombinant protein from transformed *Lactobacillus* sp. TISTR891 harboring pILK01 was higher than that of transformed lactobacilli cell harbouring pGKV8E or pILK02 in the same strain. The negative control harbouring only the cloning vector pGKV210 or pIL1020 showed no band with catalase activity (lane 2, 4, 7). Transformed *L. plantarum* TISTR850 harboring pILK01 (lane 8) showed the same pattern and intensity of catalase bands as compared to wild type and transformed *Lactobacillus* sp. TISTR891 with pILK01 (lane 1, 5). The amount of protein from transformed *L. plantarum* (lane 8) is only one third in amount as compared to the rest. Hence, the activity of transformed *L. plantarum* by in-situ activity gel assay correlates with the result obtained by quantitative catalase assay from Table 2.

#### **Nucleotide sequence of the catalase gene**

Figure 4. shows the nucleotide sequence of the 1.6 kb insert in pKP01 which encodes the catalase of *L. sake* SR911. The sequence contained 1436 bp from the structural gene of *L. sake* SR911 encoding 478 amino acids and the 17 amino acids from pMEx8 vector. The additional amino acids from cloning vector did not affect the expression of the gene. The base composition of the catalase gene has 55.41 % AT content. The nucleotide sequence shows 93.75 % homology to that of the catalase gene of *L. sake* LTH 677 (data not shown). The difference of DNA and amino acid residues between 2 strains are also indicated.

#### **Characteristics of recombinant strain *L. plantarum* TISTR 850 in culture**

Recombinant strain *L. plantarum* TISTR 850 was grown under aerobic and anaerobic conditions, and the catalase activities of intact cells and crude extracts were determined quantitatively. Catalase activity is at the highest level at stationary phase when grown under aerobic condition (data not shown). As shown in Table 3, there was no difference in the catalase activity of the transformed cells in anaerobic and aerobic

condition. The stability of the plasmid pILK01 in *L. plantarum* was stable and the expression of catalase was retained after 20 generations.

#### ***L. plantarum* TISTR 850 in fermented meat product**

The stability of plasmid under condition of meat fermentation (ie. No shaking, no adding antibiotics) was investigated. The plasmid was retained in more than 70% of the cells after 2 days of fermentation. The physical properties of fermented products using *L. plantarum* as a starter culture were similar as compared to control (ie pH, color, texture) (figure 5)

#### **Iodine number in fermented meat product**

Rancidity of the fermented meat product was determined by measuring iodine number from lipid in the products made from *L. plantarum* and recombinant *L. plantarum*. The result in Table 4 showed no difference in iodine value in lipid extracted from fermented meat product kept at 4 C for one month or at room temperature for 2, 5 and 8 days.

### **DISCUSSION**

Catalase activity in lactobacilli is a very rare property. It can prevent flavor and food color defects in fermented foods. Certain number of strains isolated from Thai fermented products were found to exhibit catalase activity if exogenous hematin was added. *Lactobacillus sake* SR911 is one of the strains that exhibits strong catalase activity and was selected for further manipulation. The main reason to transfer this gene to other catalase negative lactic acid bacteria is to improve desired characteristic of starter organisms in food fermentation to yield a more desirable product. In this study, we have successfully cloned and expressed the catalase gene of *L. sake* SR911 in *E. coli* UM2. This gene was then transferred to *Lactobacillus sp.* TISTR891 and *L. plantarum* TISTR850 which are naturally deficient in catalase.

From previous report (Noonpakdee et.al., 1996), we expressed the structural catalase gene of *L. sake* SR911 under the control of the *tac* promoter in *E. coli* UM2. However, following the subcloning of this gene into a *E. coli-lactococcus* shuttle vector pGKV210 and transforming the lactobacilli, no expression could be detected. This means that the transcription and translation signals from the strong *E. coli* promoter was probably not recognized by the lactic acid bacteria even though the so-called -35 (TTGACA) and -10 (TATAAT) boxes of

*E. coli* promoter resembles that of lactobacilli. Similar phenomenon was also described by Roy *et al.* They failed to express the manganese superoxide dismutase gene of *E. coli* in lactobacilli

(Roy *et al.*, 1993)

The 1.6 kb PCR amplified catalase gene which includes the promoter region was cloned into the vector pGKV210, and pIL1020 and expressed in both *E. coli* UM2 and lactobacilli. Both shuttle vectors pGKV210 and pIL1020 contain the same origin of replication of pWV01 derived from the cryptic plasmid of *Lactococcus lactis* subsp. *cremoris* plasmid. Plasmid pGKV210 is commonly used as a promoter screening vector but pIL1020 is an expression vector with a strong lactococcal promoter p59. The level of expression of the catalase gene under the control of its own promoter in recombinant plasmids pGKV8E and pILK02 is similar. The expression under the strong promoter p59 in recombinant plasmid pILK01, however, is approximately 2 fold higher. This is indicated by the catalase activity in transformed lactobacilli (Table 2) and by the intensity of the protein band in the region of 62,000 Da isolated from recombinant *E. coli*. The increase in the expression of catalase protein should be due to the increase in the rate of mRNA synthesis from under the strong p59 promoter or under its own promoter since transcriptional signal of p59 does not contain a Shine Dalgarno sequence (Vossen *et al.*, 1987). Brurberg *et al.* also observed higher chitinase activity under p59 control using pGKV259 to express a chitinase gene from *Serratia marcescens* in *Lactococcus lactis* and *Lactobacillus plantarum* (Brurberg *et al.*, 1994).

Interestingly, the catalase activity expressed under the control of p59 in transformed *L. plantarum* TISTR850 is increased approximately 3 fold as compared to that of *Lactococcus sp* TISTR891 and the natural host *L. sake* SR911. This explains the fact that the expression level of catalase gene under p59 in pILK01 varies among different lactobacilli species or strains.

The subunit size of the *L. sake* SR911 catalase as previously reported (Noonpakdee *et al.*, 1996) and in this study is approximately 62,000 Da which corresponds to the sizes of catalase isolated from animals, plants and microorganisms (Nadler *et al.*, 1986; Ossowski *et al.*, 1991). The catalase *in situ* activity staining in non-denaturing polyacrylamide gel shows 2 bands of proteins. A prediction as to the quaternary structure of the catalase from lactic acid bacteria may indicated that it is a hexameric and similar to catalase-1 from *B. subtilis* (Loewen *et al.*, 1987) and catalase HP11 from *E. coli* (Ossowski *et al.*, 1991). However, the conditions in the gel may cause a partial breakdown of the

catalase fusion protein product to a trimeric form which may be the smallest active form of the enzyme. This phenomenon was also observed by Knauf *et al.*

It has been shown in this study that the catalase gene from *L. sake* can be transferred to negative catalase strains such as *Lactobacillus sp.* TISTR891 and *L. plantarum* TISTR850 to express catalase with the elevated amount of activity compared to the wild type. The manipulated strains may be useful in fermentation process.

*L. plantarum* harboring pILK01 has been studied in culture. Catalase activity is at the highest level at early stationary phase. There is no difference of the catalase activity when growing cells under aerobic or anaerobic condition. This may be because p59 promoter is not regulated by oxygen. The recombinant plasmid in *L. plantarum* is stable when used as a starter culture in meat fermentation. Iodine number of the fermented meat products from recombinant *L. plantarum* shows no difference to that of control.

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**Table 1.** Bacterial strains and plasmids

Bacterial strain or plasmid	Relevant properties	Source/reference
<b>Bacterial strains</b>		
<i>E. coli</i> UM2	<i>katE2 katG15</i>	12
<i>Lactobacillus sake</i> SR911	wild type, heme catalase	11
<i>Lactobacillus sake</i> LTH677	wild type, heme catalase	6
<i>Lactobacillus</i> sp. TISTR891	catalase negative	11
<i>Lactobacillus plantarum</i> TISTR850	catalase negative	11
<b>Plasmids</b>		
pUC19	Ap <sup>r</sup> , lacZ, 2.7kb	13
pUCK	Ap <sup>r</sup> , lacZ, 3.3kb, pUC19 carrying the catalase gene <i>KatA</i>	This study
pGKV210	Em <sup>r</sup> , Cm <sup>r</sup> , 4.5 kb, pW01 replicon	14
pIL1020	Em <sup>r</sup> , Cm <sup>r</sup> , 5.1 kb, pW01 replicon	INRA
pGKV8E	Em <sup>r</sup> , Cm <sup>r</sup> , 6.1 kb, pGKV210 carrying the catalase gene <i>KatA</i>	This study
pILK01	Em <sup>r</sup> , Cm <sup>r</sup> , 6.7 kb, pIL1020 carrying the catalase gene <i>KatA</i>	This study
pILK02	Em <sup>r</sup> , Cm <sup>r</sup> , 6.7 kb, pIL1020 carrying the catalase gene in opposite direction	This study
pILK03	same as pILK02	This study

Abbreviations: TISTR, Thailand Institute of Scientific and Technological Research; INRA, Institut National de la Recherche Agronomique Research Center, Jouy-en-Josas, France; Em<sup>r</sup>, erythromycin resistance; Cm<sup>r</sup>, chloramphenicol resistance; Ap<sup>r</sup>, ampicillin resistance



**Table 2.** Catalase activity expressed by transformed lactobacilli harboring the gene encoding the catalase of *L. sake* SR911

Strains	Catalase activities	
	$\mu\text{moles of H}_2\text{O}_2 \text{ decompose} \times \text{min}^{-1} \times \text{mg of protein}^{-1}$	
<i>L. sake</i> SR 911(wild type)	154	
<i>Lactobacillus sp.</i> TISTR891	0	
<i>Lactobacillus sp.</i> (pGKV210)	0	
<i>Lactobacillus sp.</i> (pGKV8E)	67	
<i>Lactobacillus sp.</i> (pIL1020)	0	
<i>Lactobacillus sp.</i> (pILK01)	167	
<i>Lactobacillus sp.</i> (pILK02)	74	
<i>Lactobacillus sp.</i> (pILK03)	82	
<i>L. plantarum</i> TISTR850	0	
<i>L. plantarum</i> (pILK01)	492	

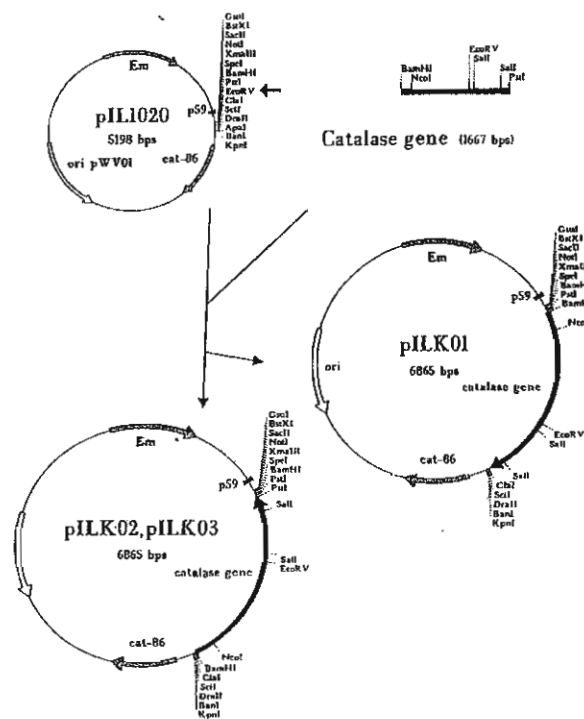
<sup>a</sup> Data represent average number of 3 experiments, each performed in duplicate.

**Table 3.** Catalase activity of *L. plantarum* and *L. plantarum* harbouring pILK01 under aerobic and anaerobic condition

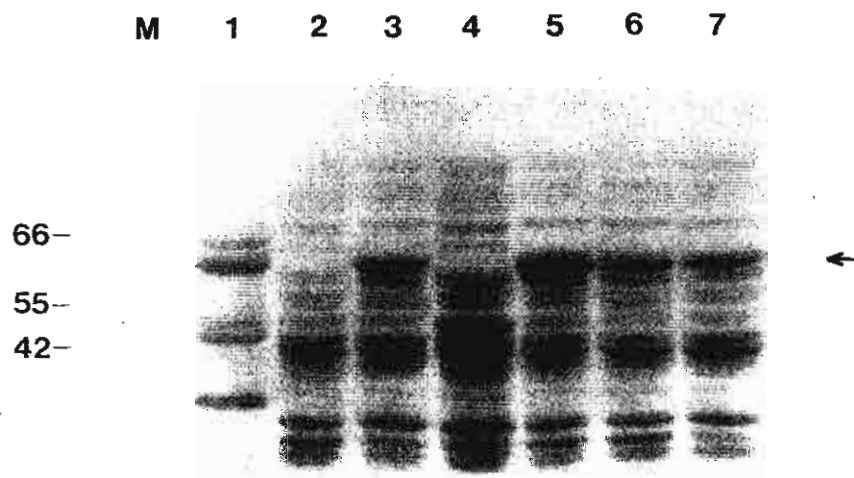
Strain	Growth condition	Catalase activities ( $\mu\text{moles H}_2\text{O}_2 \text{ decomposed/min.}$ )	
		/OD <sub>600</sub>	/mg protein
<i>L. plantarum</i>	aerobic	0	0
	anaerobic	0	0
<i>L. plantarum</i> (pILK01)	aerobic	70.3	233
	anaerobic	87.2	223

**Table 4.** Iodine number of extracted lipid from Nham prepared from *L.plantarum* TISTR850  
*L.plantarum* TISTR850 harboring pILK01

Starter	Iodine number					
	0 day	at room temperature			at 4 °C	
		2 days	5 days	8 days	1 month	2 months
Control	81.51	-	-	-	87.98	-
<i>L.plantarum</i> TISTR850	83.17±3.38 (n=10)	85.70±4.43 (n=2)	83.47±1.87 (n=3)	81.97±2.54 (n=3)	86.16±2.46 (n=5)	-
<i>L.plantarum</i> TISTR850 harboring pILK01	84.30±3.41 (n=9)	87.94±0.23 (n=3)	82.69±5.39 (n=4)	78.49±1.57 (n=4)	83.94±1.97 (n=7)	-



**Figure 1.** Schematic representation of the construction of a plasmid pILK01, pILK02, and pILK03 using *E. coli-lactococcus* expression vector pIL1020. The linear fragment represents the fragment of catalase gene including its promoter region. P59 is lactococcal promoter region. Arrows marked ori denote DNA regions required for replication and the transcriptional direction of the major open reading frames in the region. EmR and CmR indicate resistance to erythromycin and chloramphenicol respectively.



**Figure 2.** SDS-PAGE of total soluble proteins of transformed *E. coli* UM2 harboring the gene encoding the catalase gene of *L. sake* SR911. Lanes: M, molecular weight marker: 1, *L. sake* SR911; 2, *E. coli* UM2(pGKV210); 3, *E. coli* UM2 (pGKV8E); 4, *E. coli* UM2 (pIL1020); 5, *E. coli* UM2 (pILK01); 6, *E. coli* UM2 (pILK02); 7, *E. coli* UM2 (pILK03) .



**Figure 3.** Catalase activity staining in PAGE of soluble proteins of strains harboring the gene encoding the catalase gene of *L. sake* SR911. Lanes: 1, *L. sake* SR911; 2, *Lactobacillus* sp.TISTR891(pGKV210); 3, *Lactobacillus* sp. TISTR891(pGKV8E); 4, *Lactobacillus* sp.TISTR891(pIL1020); 5, *Lactobacillus* sp.TISTR891(pILK01); 6, *Lactobacillus* sp.TISTR891 (pILK02); 7, *L. plantarum* TISTR850(pIL1020); 8, *L. plantarum* TISTR850(pILK01).

**Figure 4.** Nucleotide sequence of the catalase gene and deduced amino acid.

## เนื้อหางานวิจัย

### TRANSFORMATION OF THAI PAPAYA CULTIVAR BY PARTICLE GUN BOMBARDMENT

#### ABSTRACT

Embryogenic calli and somatic embryos of Thai papaya cultivar Khakdum were transformed with binary vector containing kanamycin as a selectable maker. The tissues were bombarded by M10 tungsten and 1.0  $\mu\text{m}$  gold particles at helium pressure 1300 psi. After bombardment transgenic calli were selected on the media containing 100 mg/l kanamycin. One transgenic callus was obtained from 810 zygotic embryos of embryogenic calli bombarded with M10 tungsten particles (0.12%) and non transgenic callus was obtained from 1590 embryogenic calli bombarded with and 1.0  $\mu\text{m}$  gold particles. Somatic embryos obtained from the culture of papaya embryos grew on the media containing 10 mg/l 2,4-D for 2-3 months are more susceptible for the transformation than the embryogenic calli. Seven transgenic calli were obtained from 1200 clumps of somatic embryos bombarded with M10 tungsten particles (0.58%) and four transgenic calli from 780 clumps of somatic embryos or about 0.51% were obtained from somatic embryos. Transgenic papaya plants cultivar Khakdum were successfully regenerated and transferred to soil.

#### INTRODUCTION

Papaya (*Carica papaya* L.) is a fruit crop grown in tropical and subtropical countries. In Thailand, papaya is one of the staple food. The green fruits are used as salad and the ripened fruits are consumed as fresh dessert. Papaya trees are commercially grown in plantations and on a small scale in home gardens. Since 1975, the papaya production in Thailand has been limited by papaya ringspot virus (PRSV). The major cultivars of papaya grown in Thailand, Khak Dum and Khak Nuan are very susceptible for this virus. This virus causes seriously damage to papaya plantation. The infected area increases each year and it is now endemic throughout Thailand.

The use of genetic engineering to produce transgenic papaya by *Agrobacterium* and microprojectile bombardment had been reported (Fitch et al, 1990; Fitch et al, 1992, Cabrera-Ponce et al, 1995; Yang et al, 1996, Cheng et al, 1996). In these experiments the transgenic Hawaiian papaya cultivar containing the coat protein gene of PRSV mild strain isolated from Hawaii (HA 5-1) were generated. The transgenic papaya showed resistance to the severe strain of PRSV isolated from Hawaii (HA). However this transgenic plant was not

resistance to PRSV isolated from Thailand and other countries (Tennant *et al.*, 1994). Further the papaya cultivar. "Sunset" and "Kapoho" used in these experiments are not widely consumed in Thailand. In this research we report the DNA transfer of binary vector into Thai papaya cultivar Khakdum using microprojectile bombardment.

## **MATERIALS AND METHODS**

### **Plasmid**

The binary vector pSA1006 is based on pGV941 (Deblaere *et al.*, 1987) and contains the *nos* promoter- *nptII* gene-3'*nos* as a selectable marker and the CaMV35S promoter-*uidA* gene-3'*nos* from pBI 121 (Jefferson, 1987) as a reporter gene.

### **Plant material and culture**

Immature zygotic embryos were dissected from seeds of immature fruits of papaya var. Khakdum. Zygotic embryos were cultured on the induction medium (M1 medium) which consisted of: half strength MS salts medium (Murashige and Skoog's, 1962) plus 50 mg/l myo-inositol, full strength MS vitamins, 400 mg/l glutamine, 10 mg/l 2,4-D, 6% sucrose, 8 g/l Difco Bactoagar, pH 5.8. After the zygotic embryos were maintained in the dark at 26°C for 3-4 weeks, the embryogenic calli which produced one or more somatic embryos on the apex were developed. These embryogenic calli were used in this transformation experiment. The second type of papaya culture used in this experiment is the somatic embryos. A small clump of somatic embryos was developed on the embryogenic callus when the callus was maintained in the induction medium for two to three months with monthly subcultured. Both types of tissues were transferred to the fresh M1 medium in group of 30 calli per petri dish one week before the bombardment

### **Plant transformation**

Embryogenic calli and somatic embryos were bombarded with M10 tungsten and 1.0 µm gold particles coated with the binary vector using the PDS 1000/helium-driven biolistic device (Bio-Rad, Hercules, Calif.). The gap distance between the rupture disk and the macrocarrier is 1.2 cm and the petri dish was placed at the target level 2 (6.0 cm). The chamber is evacuated to 26 in Hg and the helium gas pressure is set at 1300 psi. After bombardment the embryogenic calli and somatic embryos were cultured in the M1 medium for 2 days and transferred to the callus propagation medium (M1 medium supplemented with 75 mg/l kanamycin and 500 mg/l cefotaxime). After 14 days the tissues were transferred to the selective media which is the callus propagation medium containing 100 mg/l kanamycin.

The tissues were subcultured monthly to fresh selective media until resistant embryogenic calli developed.

### **Plant regeneration**

The resistant embryogenic calli were transferred to glass vessels with plastic lids containing the regeneration medium (MS medium supplemented with 0.2 mg/l BAP, 0.1 mg/l kinetin, 3% sucrose, 8 g/l Difco Bacto agar, 100 mg/l kanamycin, pH5.7). The tissues were cultured under 12 hour photoperiod using cool white Sylvania fluorescent lamps until the embryos regenerated and formed shoots. Shoots with 3-4 leaves and about 1.5 cm tall were sliced and transferred to the rooting medium. Two formulas of rooting media were used in this experiment. The R1 medium is a hormone free medium containing half strength MS salts, half strength MS vitamins, 1.5% sucrose, 8 g/l Difco Bactoagar at pH 5.8. The R2 medium contains full strength MS salts, full strength MS vitamins, 100 mg/l myo-inositol, 1 mg/l indol butyric acid, 3% sucrose, 8 g/l Difco Bactoagar at pH 5.8. Rooting plants were acclimatized by replaced the plastic lids of the culture vessels with a sheet of double layer sterile tissue paper and covered with saran wrap for 2 weeks before the plant transferred to 1:1 mixture of soil and vermiculite.

### **GUS histochemical staining**

The kanamycin resistance callus was analyzed for  $\beta$ -glucuronidase activity by histochemical assay (Jefferson, 1987). About 50 mg of callus was placed in a 1.5 ml microcentrifuge tube and incubated in 100  $\mu$ l of 1 mM X-gluc in 50 mM phosphate buffer for 1-16 hr at 37°C.

## **RESULTS AND DISCUSSION**

### **Transformation of Thai papaya cultivar Khakdum**

Two type of papaya cultures were used for DNA transformation by particle gun bombardment, the zygotic embryos and the clump of somatic embryos. After bombardment the cultures were transferred to the selectable medium containing 100 mg/l kanamycin and transferred monthly to fresh selective medium until transgenic calli were selected which took about 4-5 months. Transgenic calli were identified as light-yellow somatic embryo clusters among the brown tissues of the non-transformed calli. Zygotic embryos of Thai papaya cultivar Khakdum are not susceptible for particle gun bombardment when compared with somatic embryos. After the tissues were cultured on selective medium for 5 months only one transgenic callus was obtained from 810 zygotic embryos bombarded with M10 tungsten particles (0.12%) and non of the transgenic callus was obtained from 1590 zygotic embryos

bombarded with 1.0  $\mu\text{m}$  gold particles. In the experiment using somatic embryos for bombardment seven transgenic calli were obtained from 1200 clumps of somatic embryos bombarded with M10 tungsten particles (0.58%) and four transgenic calli were obtained from 780 clumps of somatic embryos bombarded with 1.0  $\mu\text{m}$  gold particles (0.51%). The efficiency of transformation of papaya cultivar. Khakdum using somatic embryos is comparable to the previously reported in particle gun bombardment of zygotic embryos of papaya cultivars "Sunset" (Fitch *et al*, 1990) and " Maradol" (Cabrera-Ponce *et al*, 1995). However the papaya cutivar Khakdum took about 1 month longer to regenerate. The genotype of the papaya, the type and age of tissue used in the particle gun bombardment are the important factors that effect the transformation efficiency by particle gun bombardment .

### **Plant regeneration**

The transgenic calli started to regenerate after culturing for 3-4 months on the regeneration medium containing NAA, BAP and kanamycin and multiple shoots were formed within 6-9 months. Shoots with 3-4 leaves and about 1.5 cm tall were sliced and transferred to the rooting medium. When using media R2 which containing 1 mg/l indol butyric acid for rooting, few thick and short roots are formed at the end of the shoot. However this type of root is not healthy and not elongate when compared with the root of plant grew on the media R1 which is hormone free media. Thus the regenerated shoots are cultured in the R1 media until the thin long healthy roots formed which took about 1 month and then transferred to soil.

### **GUS expression in transgenic calli.**

Transgenic calli resistant to 100 mg/l kanamycin were assayed for the  $\beta$ -glucuronidase expression by histochemical test. Five transgenic lines out of eight bombarded with M10 tungsten particles and one transgenic line out of four bombarded with gold particles showed *gus* expression in form of spot staining on the callus. Some reports suggested that transgenic plants may not express the *gus* gene product due to the deletion of the *gus* gene from the plasmid which was subjected to fragmentation during particle gun bombardment and the gene might not be linked to the kanamycin resistance gene (Fitch *et al*, 1990). Moreover, it has been also reported that the papaya callus tissue might contain detrimental factors leading to degradation of the GUS transcript and the GUS protein (Yang *et al*, 1996).



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## เนื้อหาทางวิจัย

### CHARACTERIZATION OF EXPRESSED SEQUENCE TAGS FROM BLACK TIGER SHRIMP 'S HEMOCYTES

#### ABSTRACT

Two lambda ZAPII-based cDNA libraries was prepared from poly(A)+ RNA of hemocytes from Black Tiger Shrimp (*Penaeus monodon* Fabricius). Random clones were isolated and their 5' end partial nucleotide sequence determined by automated procedure. A segment of each DNA sequence was used to search online nucleotide and protein sequence databases by two BLAST programs. From the first cDNA library with a lower phage titer, 53 DNA sequences were obtained. Among these, 12 sequences (22%) did not match existing DNA sequence in online databases by either BLASTN or BLASTX searches, while 12 sequences (23%) matched known DNA sequences for known protein identities and 29 sequences (55%) matched 16S ribosomal rRNA sequences. Among those matching protein coding sequences, 3 sequences (6%) matched those of unique genes, 5 sequences (9 %) matched those of ribosomal protein genes, 4 sequences (7%) matched those of mitochondrial genes. A second cDNA library constructed from higher amount of poly(A)+ RNA provided higher phage titer. Analysis of random cDNA sequences from the second cDNA library was in progress. The study of partial cDNA sequences should allow rapid characterization of the DNA sequences, some of which may be relevant to immune system of the shrimp.

#### INTRODUCTION

Shrimps in the subphylum Crustaceae are a group of highly diversified organisms of over 38,000 known species (1). Human consume a number of marine shrimps species, yet those in the Family Penaeidae constitute the bulk of the catches (2). With the adoption of certain fast growing *Penaeus* species for aquaculture around the World, its commercial impact is even higher. In South and Southeast Asia, *Penaeus monodon* is the choice species. For Thailand alone, the World 's number one exporter of frozen shrimps, a yearly export value for about 250,000 tonnes of frozen shrimps amounted to US\$ 2 billions.

Despite its economic importance to the World, little attention has been paid to study the shrimps in Molecular Biology aspects. Online DNA sequence databases possessed only a hundred plus records, as of early 1999. In recently years' a number of viruses infecting

Penaeid shrimps have been identified which caused epidemics among the farm-reared *P. monodon* shrimp of several countries (3,4,5). Outbreak of viral diseases such as white spot disease and yellow head disease for *P. monodon* in 1996 destroyed almost half of Thailand's annual shrimp output. Sporadic infections of those viruses were still reported in many localities. It is clear that works in the DNA level on the shrimp and its viruses have become an urgent necessity.

Work on cDNA library and the clones in various organisms have allowed rapid identification of expressed sequences (6). Such partial cDNA sequences, widely known as Expressed Sequence Tags (or EST), are well characterized in human with over one million sequence records believed to cover most of the functioning human gene sequences. There are many useful application of EST information, such as serving as gene-specific tags in DNA cloning and gene characterization, as chromosomal markers for selective breeding programs and for genome mapping research, as known targets for PCR amplification in preparation of DNA probes, etc. With the severe deficiency in Penaeid shrimps DNA sequence data, as well as overall deficiency for those across the wide taxa of crustacea as well, it was also very difficult for researchers to start working on any gene systems of the shrimp although some information of the particular genes might already be available from other invertebrate models such as *Drosophila* and *Caenorhabditis*. To accumulate data fast with minimal cost, we had embarked on characterization of the shrimp's random cDNA clones from muscular tissues to prepare ground work on the shrimp's molecular genetics (7). Result from EST work had let our group to identify and characterize full-length sequences of many cDNA from the black tiger prawn, such as enolase (8), two actin sequences (9), arginine kinase (10).

With some known cDNA sequences serving as our foundation for addressing other molecular biology questions, we then started to ask question on how much did we know about on the genes relevant to the defense and immune mechanism of the shrimp. Limited information from work conducted in a primitive crustacean, *Limulus polyphemus*, and many insect species which belong to the same phylum indicated that phagocytosis is the main defense mechanism (11). Hemocyte became the natural choice for us as starting material for collecting gene information by the same approach. In this communication, we describe our recent work on the EST from *P. monodon*'s hemocytes.

## MATERIALS AND METHODS

### Preparation of cDNA library and isolation of cDNA clones

Two months old *Penaeus monodon* shrimp specimens (hatched from eggs of a mother prawn caught from Andaman sea) were purchased from local farms in western Thailand. Live shrimps were briefly rinsed in sterile distilled water, then the hemolymph was quickly withdrawn from the cardiac by a sterile 22 G needle and chilled in ice until sufficient volume was obtained. The first preparation of the total RNA for preparation of the first cDNA library was prepared directly from whole hemolymph by extracting by Trizol reagent. The second cDNA library was prepared from pelleted hemocytes by the acid guanidiniethiocyanate method (12,13). Poly (A)+ RNA was obtained by means of either an oligo(dT) cellulose column (14) or magnetic procedure. About 0.15 ug and 2 ug of poly (A)+ RNA were used in the two library constructions, respectively. A cDNA synthesis kit of Strategene was used and the cDNA was cloned into EcoRI - XhoI site of a lambdaZAPII in *E.coli* XL1-Blue MRF' host, using Gigapack III Gold *in vitro* packaging kit. The two cDNA libraries yielded unamplified titers of around  $4 \times 10^4$  pfu and  $3 \times 10^5$  pfu, respectively. An aliquot from the unamplified library was taken and plated at one time to give few hundred pBluescript SK' phagemid clones using supertransfection of a f1 helper (ExAssist) phage. Random clones were picked from white colonies in LB medium containing X-gal and IPTG, and separately grown at 37 C in LB medium containing 50 ug / ml Ampicillin for 18 hr. Plasmids were extracted and partially purified by CTAB procedure (14) and cleaned up by QIAGEN. Clones whose inserts could be recovered by cutting with Eco RI and Xho I were selected for sequencing.

### DNA sequence determination

Each plasmid DNA was sequenced (15) by using AmpliTaq DNA Polymerase FS enzyme and a sequencing kit from Perkin-Elmer. Reactions were performed mainly with the M13 Universal reverse sequencing primer in a thermal cycler (Perkin Elmer model 2400) set at 10 sec at 96 C, 5 sec 50 C, and 4 min 60 C for 25 cycles. The reactions were then fractionated and analyzed by Perkin Elmer ABI Prism model 377 automated DNA sequencer, using 6% acrylamide (+ 7 M Urea) running xxx volts, xxx watts, for about 10 hr.

### DNA sequence search and alignment

DNA sequences from the regions of clear chromatogram (typically 400-500 bp out of a maximum sequence read from the machine) were used in the computer searches of GenBank (16), conducted via the world-wide web server of NCBI (<http://www.ncbi.nlm.nih.gov>).

The program used was BLASTN 1.4.9 MP and BLASTX ( 17, 18). The cDNA clones which showed statistically significant matching scores were then individually compared to some selected DNA sequences by ClustalW program, preferentially to *Drosophila melanogaster*, if available. Most of 5'-EST sequences reported in this paper will be deposited in the dbEST database. Only few upon which complete sequences were later determined and confirmed will be reported elsewhere.

## Result and Discussion

A first *P.monodon* 's hemocyte cDNA library was constructed from a small amount of poly(A)+ RNA and was found to have a low unamplified titer of around  $4 \times 10^4$ . The insert sizes of most clones range from 0.5 kb upto about 1 kb. Few hundred clones were isolated and verified that each possessed an insert cloned in between correct restriction sites (EcoRI and Xho I) as used in the library construction procedure before they were chosen to be excised in the form of pBluescript SK- phagemid. They were later sequenced using fluorescent dye-terminator chemistry on a Perkin-Elmer ABI Prism 377. Screening of several hundred clones revealed a small fraction, 53 clones which could be digested with the restriction enzymes and thus considered as containing legitimate inserts. Most clones gave DNA sequences, with the readable portion was about 300-500 bases for each clone. When only the partial sequence from each clone was used for sequence query via the Blast programs, 12 sequences (22% of 53 clones) did not match known gene sequences in existing databases by either BLASTN or BLASTX searches. This might that the shrimp DNA sequences were quite diverged from those of known eukaryotic species including those of *S.cerevisiae* and *C.elegans*. Only 12 sequences (23%) matched known gene sequences and 29 sequences (55%) matched 16S ribosomal rRNA sequences. Among those matching protein coding sequences, 3 sequences (6%) matched those of unique genes, 5 sequences (9 %) matched ribosomal protein gene sequences, 4 (7%) sequences matched mitochondrial genes.

Due to high level of contamination of the putative 16S rRNA sequences in the cDNA library, a decision was made to prepare a second cDNA library using much more hemolymph sample. The protocol was also slightly modified such that total RNA was extracted from pelleted hemocytes (which include phagocytic cells) and thus higher amount of poly(A)+ RNA was obtained. That resulted in much higher titer of the recombinant lambda phage of around  $3 \times 10^5$  pfu. Analysis of random cDNA sequences from a second cDNA library was still in progress.

The availability of our DNA sequences in the online databanks should facilitate investigators working in other invertebrates in identifying their gene products of known or unknown function in the near future. The study of partial cDNA sequences allow rapid characterization of the DNA sequences some of which may be relevant to immune system of the shrimp.

**Table I** List of putative identity of some *P.monodon* EST from the first hemocyte cDNA library.

Clone ID	Putative Identity	Ref. organism (with accession no.)	sequence% identity
PMH08	Thymosin beta-11	(CAA21832)	38 %
PMH17	Neuronal protein NP45 or Calponin homolog	Rattus norvegicus (M84725) Schistosoma mansoni (AAB47536)	91% 52 %
PMH25	ADP-ribosylating factor	A. thaliana (M95166)	88 %
PMH38	Ribosomal protein S12	Sus scrofa (P46405)	79 %
PMH74	Cytochrome oxidase subunit 1	Penaeus northalis (CAB40364)	75 %
PMH83	NADH dehydrogenase subunit 5,	Panorpa japonica (AF056494)	83 %
PMH95	Ribosomal protein S5	S.pombe (AB029515)	83 %
PMH140	Ribosomal protein P2, EL12	Artemia sp. (P02399)	39 %
PMH172	Cytochrome oxidase III	Penaeus northalis (X84350.2)	85 %

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## เนื้อหางานวิจัย

### MOLECULAR CLONING OF TWO CDNAS ENCODING NEUROPEPTIDES IN CHH/MIH/GIH FAMILY FROM *PENAEUS MONODON* AND THEIR GENE STRUCTURE

#### ABSTRACT

Crustacean hyperglycemic hormone (CHH), molt-inhibiting hormone (MIH) and gonad-inhibiting hormone (GIH) are crustacean neuropeptides that control sugar metabolism, molting process and gonad maturation, respectively. They are synthesized by the X-organ-sinus gland complex (XOSG) in the eyestalks. In this study, we used a technique of Rapid Amplification of cDNA Ends (RACE) to clone cDNAs that encode peptide hormones in CHH/MIH/GIH family from *Penaeus monodon*'s eyestalks. Two cDNA clones, *Pem-CMG1* and *Pem-CMG2*, were obtained. *Pem-CMG1* and *Pem-CMG2* coded for putative peptides of 127 and 128 amino acids, respectively. The putative mature peptides of both *Pem-CMG1* and *Pem-CMG2* comprised 74 amino acid residues. These two peptides shared 43% identity in amino acid sequences of the leading peptide region whereas 95% identity was found in their mature peptide sequences. Genomic fragments of *Pem-CMG1* and *Pem-CMG2* were obtained by PCR. The gene for *Pem-CMG1* was composed of two exons that were separated by an intron of 314 nucleotide pairs in the mature peptide coding region. *Pem-CMG2* gene harbored two introns, one was 405 nucleotides in length inserting into leading peptide coding region. The other intron was 268 nucleotides long inserting into a coding region for mature peptide. These results suggest that *Pem-CMG1* and *Pem-CMG2* were encoded by two related genes that have similar sequences in the coding region for their mature peptides but are structurally different.

#### INTRODUCTION

A number of physiological processes in decapod crustaceans are known to be regulated by diverse neuropeptides synthesized by a neurosecretory system called X-organ-sinus gland (XOSG) complex located in the optic ganglia of the eyestalks (1). In contrast to most of the peptide hormones isolated from the XOSG complex e.g. pigment dispersing hormone (PDH) and crustacean cardioactive peptide (CCAP) that have also been identified in insect, a group of related peptides constituting a novel hormone family has been described and has thus far been found only in crustaceans (2). This family includes crustacean hyperglycemic hormone (CHH), molt-inhibiting hormone (MIH) and gonad

inhibiting hormone (GIH) that are involved in blood sugar regulation, inhibition of ecdysteroid synthesis and regulation of reproduction, respectively (3, 4).

Amino acid sequences of the hormones in this CHH/MIH/GIH family have recently been determined in several crustaceans (5-8). Comparison of amino acid sequences of these hormones revealed significant degrees of similarity (2, 9) suggesting that these hormones constitute an authentic peptide family across species boundaries. In addition to several similarities in their structure, some hormones in the CHH/MIH/GIH family also exhibit overlapping biological activities (7, 10, 11). However, the certainty concerning the primary physiological functions of different peptides still needs to be clarified.

In recent years, the cDNA coding for CHH, MIH and GIH have been cloned based on the information of their available amino acid sequences (8-14). Despite this increasing number of reports on cDNA cloning and nucleotide sequence analysis of neuropeptides in the CHH/MIH/GIH family, the hormones in several crustaceans still await complete characterization, especially at molecular level.

We report here the molecular cloning of two cDNAs coding for member of CHH/MIH/GIH family, Pem-CMG1 and Pem-CMG2, from black tiger prawn, *Penaeus monodon*. Open reading frames of *Pem-CMG1* and *Pem-CMG2* cDNAs were cloned and primary structure of the genes encoding these two peptides were also analyzed.

## **MATERIAL AND METHODS**

### **Animals**

Black tiger prawns, *Penaeus monodon*, were obtained from Shrimp Culture Research Center, Chareon Pokphand Group and were kept in aerated seawater at room temperature until used.

### **Oligonucleotide primers**

Abridged anchor primer (AAP) and abridged universal amplification primer (AUAP) were provided with the 5'RACE System for Rapid Amplification of cDNA Ends (GIBCO BRL). Other oligonucleotides were purchased from Bio-Synthesis, USA. The sequences of the primers are shown in Figures 1 and 2.

### **RNA Isolation**

A total of 100 eyestalks were collected from *Penaeus monodon* immediately after anesthetized with crushed ice. After the cuticle and non-neural tissues were removed, the dissected eyestalks were ground to fine powder in liquid nitrogen by means of mortar and pestle. Total RNA was extracted using TRIzol Reagent (GIBCO BRL). Poly (A)<sup>+</sup> RNA was

isolated using Dynabeads mRNA purification kit (DYNAL) and was quantified by measuring the absorbancy at 260 nm.

#### **Amplification of the 3' ends of cDNA**

The reaction was carried out according to the protocol of the 3' RACE System for Rapid Amplification of cDNA Ends (GIBCO BRL). First strand cDNA was synthesized from 75 ng of poly (A)<sup>+</sup> RNA in a total volume of 20 µl containing 20 mM Tris-HCl (pH 8.4), 50 mM KCl, 2.5 mM MgCl<sub>2</sub>, 10 mM DTT, 500 nM PRT primer, 500 mM each dATP, dCTP, dGTP, dTTP and 200 units of SuperScript II Reverse Transcriptase. The reaction was incubated at 50°C for 50 minutes and then terminated at 70°C for 15 minutes. The RNA template was degraded with RNaseH before proceeding to amplification step. A 2µl aliquot of cDNA was amplified with primers PM1 and PM2 (see Figure 1A) in a reaction containing 20 mM Tris-HCl (pH 8.4), 50 mM KCl, 2 mM MgCl<sub>2</sub>, 200 nM each primer, 200 mM each dATP, dCTP, dGTP, dTTP and 2.5 units of AmpliTaq DNA polymerase (Perkin Elmer). Amplification was performed in a DNA Thermal Cycler (GeneAmp System 2400, Perkin Elmer) with 35 cycles of 94°C for 1 minute, 50 °C for 1 minute and 72°C for 2 minutes following with 7 minutes incubation at 72°C as a final extension.

#### **Amplification of the 5' ends of cDNA**

The detailed protocol of the 5' RACE System for Rapid Amplification of cDNA Ends Version 2.0 (GIBCO BRL) was followed. First strand cDNA synthesis was carried out as described for the 3' RACE except that 100 nM of RM3 primer was substituted for PRT. The cDNA synthesized was then purified using a GlassMax DNA isolation spin cartridge (GIBCO BRL).

A 10 µl aliquot of purified cDNA was tailed with dCTP in 25 µl of 10 mM Tris-HCl (pH 8.4), 25 mM KCl, 1.5 mM MgCl<sub>2</sub>, 200 mM dCTP and 1 µl of terminal deoxynucleotidyl transferase (TdT). The reaction was incubated at 37°C for 10 minutes and TdT was heat inactivated at 65°C for 10 minutes. The PCR amplification using 5 µl of dC-tailed cDNA was performed as described for 3' RACE using 200 mM of PM4 and abridged anchor primer (AAP). The second round PCR was performed using primer PM5 and abridged universal amplification primer (AUAP) to obtain specific amplified product (Figure 1).

#### **Amplification of an open reading frame of Pem-CMG1 peptide**

Total RNA extracted from eyestalks and primer CMG-R were used to synthesize a cDNA template for amplification of the open reading frame of Pem-CMG peptide. The

condition for PCR was the same as described earlier. The primers used were CDF and CDR, the sequences of which are given in figure 1.

#### **Amplification of an open reading frame of Pem-CMG2 peptide**

A cDNA synthesized with PRT primer was used as a template for amplification of the open reading frame of Pem-CMG2 peptide. The condition for PCR was the same as described earlier. The primers used were CDF-2 and PM1, the sequences of which are given in figure 1.

#### **Amplification of a genomic fragment of *Pem-CMG* genes**

Genomic DNA was prepared from abdominal muscle tissues of *P. monodon* by phenol/chloroform extraction method (15). About 150 ng of genomic DNA was used as a template for PCR amplification of the *Pem-CMG* sequences with the primers CMG-F and CMG-R for *Pem-CMG1* and primers CMG-F and CMG-R2 for *Pem-CMG2* (figure 2). The PCR reaction mixture (150 ng of genomic DNA template, 20 mM Tris-HCl (pH 8.4), 50 mM KCl, 2 mM MgCl<sub>2</sub>, 200 nM each primer, 200 mM each dATP, dCTP, dGTP, dTTP) was heated to 95°C for 5 minutes then, 2.5 units of AmpliTaq was added. Amplification was achieved by 35 successive cycles of denaturation at 94°C for 1 minute, annealing at 50°C for 1 minute and extension at 72°C for 2 minutes, followed by a 10 minute final extension at 72°C.

#### **Cloning and Sequencing**

PCR Amplification products were purified using GeneClean II kit (Bio101). The purified DNA fragments were digested with appropriate restriction before ligated to pUC18 vector predigested with corresponding enzymes. The ligation products were used to transformed *E. coli* JM109 and the recombinants were screened by restriction enzyme digestion. The nucleotide sequences of the recombinants were determined by the method of ABI PRISM™ Dye Terminator Cycle using ABI PRISM™ Model 377 automated DNA Sequencer (PE Applied Biosystems).

### **RESULTS**

#### **3'RACE cloning of cDNAs encoding CHH/MIH/GIH peptides**

Five different 3' fragments that shared homology in nucleotide sequences to those of CHH/MIH/GIH from other crustaceans were obtained. Nucleotide sequence comparison showed that these 3'RACE clones shared 53 % identity in nucleotide sequences of their

coding regions whereas much lower level of identity was observed among their 3' untranslated regions (figure 3).

#### **5' RACE cloning of cDNA encoding Pem-CMG peptide**

Using three specific primers generated from the nucleotide sequence of the 3' RACE-5 fragment, a single band of amplification product of 370 bp (excluding G-rich tail) was obtained from 5' RACE and was subsequently cloned into pUC18. The nucleotide sequence analysis revealed that it contained cDNA insert homologous to the mRNA of CHH/MIH/GIH peptide family. However, there were differences in nucleotide at 4 positions in the region that the 5'RACE and the 3'RACE-5 fragment overlapped.

#### **Amplification of an open reading frame of Pem-CMG1 peptide**

An eyestalk cDNA was synthesized with primer CMG-R that was designed from nucleotide sequences at the 3' end of the 3'RACE-5 fragment. This cDNA was used as a template for PCR with primer PM2 and CMG-R to amplify the 3' region of *Pem-CMG1* cDNA. The 5' region of *Pem-CMG1* cDNA was generated by PCR with PM3 and CMG-F, a primer that was designed from the sequence at the 5' end of the 5'RACE fragment (figure 1). Nucleotide sequences of the 3'*Pem-CMG1* fragment was identical to that of the 3'RACE-5 fragment whereas nucleotide sequences of the 5'*Pem-CMG1* was different from that of the 5'RACE fragment. An open reading frame for *Pem-CMG1* was cloned by amplification of the cDNA with primers CDF and CDR that contained the sequences from start codon and stop codon of *Pem-CMG1* (figure 1). The amino acid sequence deduced from *Pem-CMG1* is shown in figure 4.

#### **Amplification of an open reading frame of Pem-CMG2 peptide**

Oligonucleotide primer CDF-2 was designed from the start codon of 5'RACE fragment. The cDNA that was synthesized with PRT primer was amplified with CDF-2 and PM1 to obtain open reading frame as well as 3' untranslated region of *Pem-CMG2*. The deduced amino acid sequence of *Pem-CMG2* is shown in figure 4.

#### **Comparison of the amino acid sequences of Pem-CMG1 and Pem-CMG2**

The deduced amino acid sequences of Pem-CMG1 and Pem-CMG2 were compared in figure 4. They were 44 % identical in the leading peptide region whereas higher level of identity (92%) was found in their mature peptides (figure 4). This result suggests that the two *Pem-CMG* cDNAs were transcribed from two different genes in CHH/MIH/GIH family. Comparison of the amino acid sequences of the putative mature Pem-CMG1 and Pem-CMG2 to the corresponding regions of CHH, MIH and GIH from other crustaceans revealed

that Pem-CMG1 was 43, 35, 32 and 31% identical to CHH from *P. japonicus*, MIH from *P. japonicus*, MIH from *Carcinus maenas* and GIH from *Homarus americanus*, respectively. Similar levels of identity could also be found between Pem-CMG2 and the other four peptides (figure 5).

### Primary structure of Pem-CMG genes

Genomic fragment for *Pem-CMG1* and *Pem-CMG2*, designated as *g-CMG1* and *g-CMG2*, were amplified from genomic DNA of *P. monodon* with primers CMG-F and CMG-R and primers CMG-F and CMG-R2, respectively. These two genomic fragments contain nucleotide sequences from the first start codon (ATG) to the stop codon (TAG) of *Pem-CMG1* and *Pem-CMG2* genes.

Analysis of the nucleotide sequence of *g-CMG1* fragment cloned in pUC18 revealed that this fragment contained one intron that was 314 bp long. This intron separated the two exons in the coding region for the mature Pem-CMG peptide between the second and the third codons of <sup>94</sup>Arg (Figure 6). In contrast, *Pem-CMG2* fragment contained two introns. Intron 1, 405 bp long, interrupted the coding region for the leading peptide between <sup>6</sup>Leu and <sup>7</sup>Val, the second intron of 268 bp separated the mature peptide coding region between the second and the third codons of <sup>94</sup>Arg (figure 7). An exon-intron boundary was detected using Gene Finder program of Baylor College of Medicine (BCM) on the WWW. The splice junctions conformed to the splice donor and acceptor consensus sequence (16).

## DISCUSSION

The 3' and 5' RACE strategy (17) was employed in order to clone a *Pem-CMG* cDNAs due to the lack of information on amino acid sequences of the peptides in CHH/MIH/GIH family in *P. monodon*. This technique requires only a single oligonucleotide primer which, in our case, was generated from the conserved amino acid sequence CEDCYN that is found conserved among the peptides in CHH/MIH/GIH family from a number of crustacean (1)

The open reading frame of two Pem-CMG peptides, Pem-CMG1 and Pem-CMG2, from *P. monodon* was successfully cloned. The deduced amino acid sequences from both *Pem-CMG1* and *Pem-CMG2* cDNAs reveal putative mature peptides comprising 74 amino acid residues. The Pem-CMG peptides of *P. monodon* shared the high degree of identity with CHH of *P. japonicus* whereas lower degrees of identity are found between Pem-CMG sequences and MIH reported for *P. japonicus* and *C. maenas*. The GIH of *H. americanus* also shows comparable degree of identity with both Pem-CMGs of *P. monodon*.

The gene for Pem-CMG1 of *P. monodon* (*g-CMG1*) contains only one intron that separates the coding sequence for the mature peptide whereas the gene encoding Pem-CMG2 (*g-CMG2*) contains two introns, one separates the leading peptide coding sequences, the other interrupts coding sequence for the mature peptide. The structure of *Pem-CMG2* gene is similar to the primary structure of the genes encoding MIH in *Charybdis feriatius* and CHH-like peptide in *Metapenaeus ensis* that have recently been characterized (18,19). The *MIH* gene of *C. feriatius* contains two introns, intron 1 interrupts the sequence coding for signal peptide and the second intron separates the mature peptide sequence. Two introns were also found in *CHH*-like gene of *M. ensis*. They interrupt homologous regions to those of *C. feriatius* *MIH*, one in the signal peptide sequence and the other in the mature peptide sequence. The differences in the structure of genes in the same family among different species is not uncommon and may reflect their evolutionary pathway. It is, however, too early to draw any conclusion concerning gene structure of the member of CHH/MIH/GIH family because only little information has been obtained from a few species so far. The structure of the genes coding for other neuropeptides in CHH/MIH/GIH family from *P. monodon* as well as from other crustaceans needs to be explored. This would give a clue to answer the question of how these structurally different genes have evolved from related gene family.

The sequence information from several members of CHH/MIH/GIH family together with their biological activities suggest that the neuroendocrine regulation of related physiological activities in crustaceans is more complicated than previously thought. A complete understanding of these regulatory processes requires further information from molecular studies as well as physiologically studies to define more clearly the roles of individual peptide in this family.

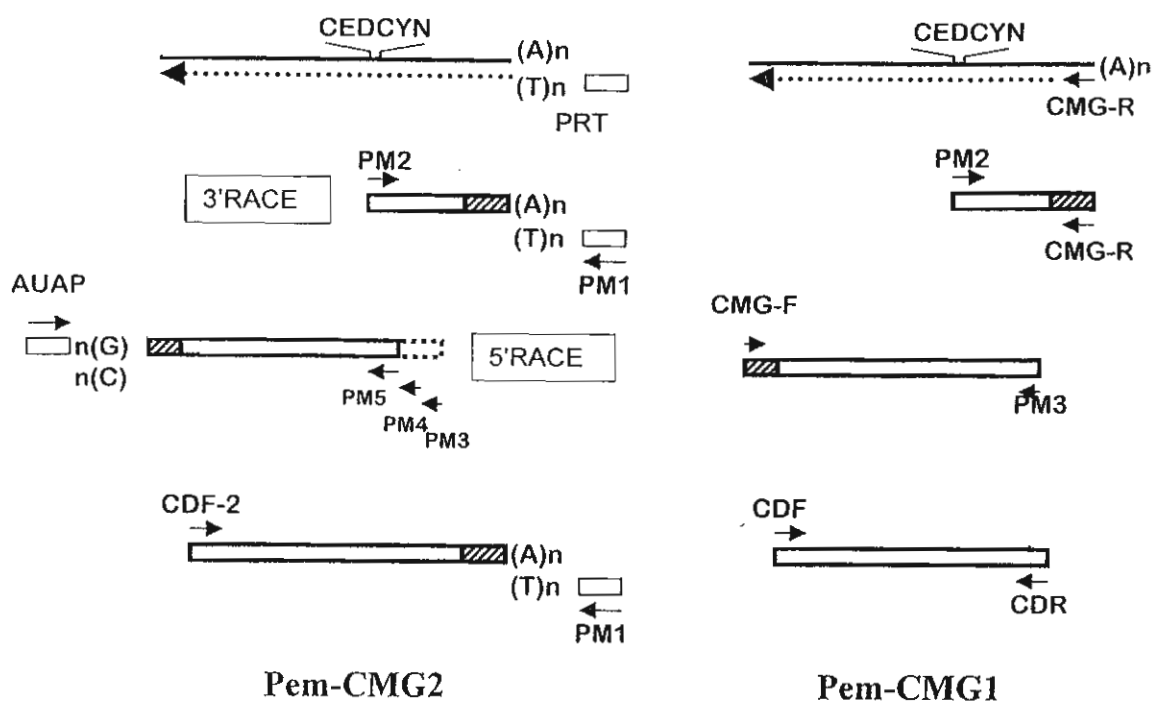
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**Figure 1.** Schematic diagram showing the strategies used for cloning *Pem-CMG1* and *Pem-CMG2* cDNA and the primers used in cDNA synthesis and PCR. Hatched boxes represent 5' and 3' untranslated regions. Detail of the primers used is given below.

#### 3'RACE primers

PRT : 5' – CCGGAATTCAAGCTTCTAGAGGATCCTTTTTTTTTTTTTTTT – 3'

PM1 : 5' – CCGGAATTCAAGCTTCTAGAGGATCC – 3'

PM2 : 5' – CCGGAATTCTGYGAAGAYTGYTACAAC – 3'

#### 5'RACE primers

AAP : 5' – GGCCACGCGTCGACTAGTACGGGIIIGGGIIGGGIIG – 3'

AUAP : 5' – GGCCACGCGTCGACTAGTAC – 3'

PM3 : 5' – CCGGAATTCCCTTTGACGAGGCCGGAAC – 3'

PM4 : 5' – CCGAAGCTTGTCCACGCAGTAGAG – 3'

PM5 : 5' – CCGGAATTCACCTCGTTGTGGAAACAG – 3'

#### open reading frame

CDF : 5' – CCGAAGCTTCATATGGTTGCCGTTGGAC – 3'

CDF-2 : 5' – CGGGATCCATGGTTGCGGTCCGATTGG – 3'

CDR : 5' – CGGGATCCCTACTTGCCGAGCCTCTA – 3'



**Figure 2.** Schematic diagram showing the strategy used for amplification of *Pem-CMG* genomic fragments and the primers used. Dotted area represent 5' and 3' untranslated regions. Primers CMG-F and CMG-R were used to amplify g-CMG1 fragment. Primers CMG-F and CMG-R2 were used for amplification of g-CMG2 fragment. Detail of the primers used is given below.

Oligonucleotide primers

CMG-F : 5' – CGGAATTCTCAGTGCAGAGGGAGAGCC – 3'

CMG-R : 5' – CGGAATTCATTTCTCTTATTTTTATTAGACAGG – 3'

CMG-R2 : 5' – GCGGATCCCTGCTTTATGAAGACACTG – 3'

```

      *      20      *      40      *      60
3'RACE-1 : TTTTCGCGTSCCAAATTTTCACCTGGTTCATTAACCTCACTCA : 61
3'RACE-2 : CTCCTALCTCGTTGGCTTGGCTTTCGACGACCAAACTTTTCAAAAGATCT : 61
3'RACE-3 : TTTTCGCGTSCCAAATTTTCACCTGGTTCATTAACCTCACTCACT : 61
3'RACE-4 : TTTTCGCGTSCCAAATTTTCACCTGGTTCATTAACCTCACTCACT : 61
3'RACE-5 : CTCCTALCTCGTTGGCTTGGCTTTCGACGACCAAACTTTTCAAAAGATCT : 61

      *      80      *      100     *      120
3'RACE-1 : CTTGGCACTGGTTCGATTACTTTCCTGGACCCAACTTCAGGAGGACATGAGGGCT : 122
3'RACE-2 : TTCTGTCTTGGTTCGATTACTTTCCTGGACCCAACTTCAGGAGGAGATGAGGGCT : 122
3'RACE-3 : TTTGGCACTGGTTCGATTACTTTCCTGGACCCAACTTCAGGAGGAGATGAGGGCT : 122
3'RACE-4 : TTCTGTCTTGGTTCGATTACTTTCCTGGACCCAACTTCAGGAGGAGATGAGGGCT : 122
3'RACE-5 : CTTGTCTTGGTTCGATTACTTTCCTGGACCCAACTTCAGGAGGAGATGAGGGCT : 122

      *      140     *      160     *      180
3'RACE-1 : GCAGACCGCTTCACTTAAGCATTTCCTTAACCTGCTTCGAGCTGGTGTGCA : 183
3'RACE-2 : GCAGAGGGCTTCACTTAAGCATTTCCTTAACCTGCTTCGAGCTGGTGTGCA : 175
3'RACE-3 : GCAGATTCGCTTCACTTAAGCATTTCCTTAACCTGCTTCGAGCTGGTGTGCA : 175
3'RACE-4 : GCAGATTCGCTTCACTTAAGCATTTCCTTAACCTGCTTCGAGCTGGTGTGCA : 175
3'RACE-5 : GCAGAGGGCTTCACTTAAGCATTTCCTTAACCTGCTTCGAGCTGGTGTGCA : 164

      *      200     *      220     *      240
3'RACE-1 : GGCTGGCTTCACCAACGGCTTCAACCACTTCCTTTTTCGCTTCGAGCTGGTGTGCA : 244
3'RACE-2 : GGCTGGCTTCACCAACGGCTTCAACCACTTCCTTTTTCGCTTCGAGCTGGTGTGCA : 235
3'RACE-3 : TTAATGCTACCTTCAGTGGACGAGAGAGGGTTTTTCGCTTCGAGCTGGTGTGCA : 236
3'RACE-4 : TCAATGCTACCTTCAGAGAGAGAGAGGGTTTTTCGCTTCGAGCTGGTGTGCA : 236
3'RACE-5 : TTTTCGCTTCGAGCTGGTGTGCATTTTCGCTTCGAGCTGGTGTGCA : 215

      *      260     *      280     *      300
3'RACE-1 : CTCGGACACCGCTTACTTTGACCTGATTCCTGCTTCGCTTCGAGCTGGTGTGCA : 304
3'RACE-2 : CTAAGTACAGCTGACCCCATATGACGCGCTGGCTTCATCTGTATGTCTTCATATA : 296
3'RACE-3 : ATTTTACCTGCTTCTGGGATTCGATATCTTTTATACCTAGATGTGTGCAAGCC : 296
3'RACE-4 : ATTTTACCTGCTTCTGGGATTCGATATCTTTTATACCTAGATGTGTGCAAGCC : 296
3'RACE-5 : CTGATTAAAGGTAATTAAGAGCCTGACACCAAGCTCTCTCTATATAAAATAAGAG : 276

      *      320     *      340     *      360
3'RACE-1 : GATCTGCTCTGTTGCTGGAGGAGGAGATGGCGTTCATGCGTCTGGTTTGGGTTGTGTCA : 365
3'RACE-2 : GCAT----- : 300
3'RACE-3 : ACTTAATGAAGTATGTTGAATT----- : 319
3'RACE-4 : ACTTAATGAAGTATGTTGAATT----- : 319
3'RACE-5 : AAATAT----- : 282

      *      380     *      400
3'RACE-1 : ATGCCTTTTCGACTCCAATAAACTTTTCAAACCTTT : 400
3'RACE-2 : ----- : -
3'RACE-3 : ----- : -
3'RACE-4 : ----- : -
3'RACE-5 : ----- : -

```

**Figure 3.** Comparison of nucleotide sequences among 3'RACE fragments. The sequence of PM2 primer is excluded from the 5' end of all fragments. The positions where the nucleotides in all five sequences are identical are highlighted (black). Dark grey and light grey area represent the positions where nucleotides are found identical in four and three sequences, respectively.

```

      *      20      *      40      *      60
CMG1 : MVAVGPMRTAVLVSLLEIPASATTFGDGNDIPTFLRSSPEASPVTSLSHTSDKRSLSFRSCTGAYD : 66
CMG2 : MVAVRLVQSAYLVSLLEIPACVTTSENTNEIPASILSSP-GDSLSEDSQISIKRSIWFNSCTGVYD : 65

      *      80      *      100      *      120
CMG1 : RELLVRLDRVCEDCYNLYRDVGVAACRSNCFHNEVFLYCVDMFRPRQRNQYRAALQRLGK : 127
CMG2 : RELLVRLDRVCEDCYNLYRDVGVAACRSNCFHNEVFLYCVDMFRPRQRNQYRAALQRLGK : 128

```

**Figure 4.** Comparison of the amino acid sequences deduced from *Pem*-CMG1 and *Pem*-CMG2. The identical amino acid residues between the two sequences are highlighted

```

      *      20      *      40      *      60
Pem-CMG1 : --SLSFRSCTGAY--DRELLVRLDRVCEDCYNLYRDVGVAACRSNCFHNEVFLYCVDMFRPR : 61
Pem-CMG2 : --SIWFNSCTGVY--DRELLVRLDRVCEDCYNLYRDVGVAACRSNCFHNEVFLYCVDMFRPR : 61
Pej-CHH : --SLFDPACTGIY--DRQLLRKLGRLCDDCYNVEREPEKVATGCRSNCFYHNLIFLDCLEYLIPI : 6
Pej-MIH : --SFIDNTCRGVMGNRDYNKKVVRVCEDCNIEFLGLDGMCRNRCFYNEWELICLKAANRED : 6
Cam-MIH : --RVINDECPNLIGNRDLYKKVEWICEDCSNIFRKTGMASLCERNCFEWEDEFVWCVHATERSE : 6
Hoa-GIH : ASAWFNDCCPGVMGNRDLYEKVAWVCNDGANIFRNNDVGVMCKKDCFHTMWELWCYATERHG : 61

      *      80
Pem-CMG1 : QRNQYRAALQRLGK--- : 74
Pem-CMG2 : QRNQYRAALQRLGK--- : 74
Pej-CHH : LQEEHMAAMQTV----- : 72
Pej-MIH : EIEKFFVWISILNAGQ- : 77
Cam-MIH : ELRDLEEWWVGILGAGRD : 78
Hoa-GIH : EIDQFFKQVSIIR----- : 76

```

**Figure 5.** Amino acid sequence alignment among the peptides in CHH/MIH/GIH family. The positions where the amino acids in Pem-CMG1 or Pem-CMG2 are identical to those of any other sequences are highlighted.

```

1   : TCAGTGCAGAGGGAGAGCCTGGAAGTTGCTGACCGTCGCTCCCGATCTGC : 50
51  : CTCTACTCTAAAGATATGGTTGCCGTTGGACCGATGCGGGCAGCTGTCCT : 100
101 : GGTGTCCCTGCTGTTGGCAATCCCGGCCTCTGCCACCACCTTCGGAGACG : 150
151 : GAAATGACATTCCAACGTTCCCTCCGTTCTTCCCCAGAAGCCTCTCCTGTG : 200
201 : ACTTCCCTTCACACCTCAGACAAACGCAGCCTATCCTTCAGGTCTTGAC : 250
251 : GGGCGCCTACGACCGCGAACTCCTTGTAAGGCTCGACCGCGTGTGCGAAG : 300
301 : ACTGCTACAACGTGTACCGCGACGTCGGAGTGGCAGCCGAATGCAGGTAA : 350
      -----
351 : CTTATTACTTTGCAGTAACCCACCCAGTTGTGTTGTGATTAAAGACTATT : 400
      ----- Intron -----
401 : GTAGAAGCGTATTAGTATAACCATCTATTACTTTAGTATATCATCTATTAC : 450
      ----- Intron -----
451 : TATCATCTATCGGACAATGATTCATACTGAGTTTGTTATCCATGGAGCT : 500
      ----- Intron -----
501 : TTCGTTAACATCGCCTGTTGCCTAAAAAGCCTCAATAATTATAAAATGTA : 550
      ----- Intron -----
551 : TAAACAGAAATATACTTTATTCCTTATATCTCTTTATTCACATAGTTATTT : 600
      ----- Intron -----
601 : TTTACATTTTCTCCTGTACGACCTACCAGTGGAACCTACATAACATTATTC : 650
      ----- Intron -----
651 : CTTCTTCAGGAGTAACTGTTTCCACAACGAGGTGTTCTCTACTGTGTG : 700
      -----
701 : GACTACATGTTCCGGCCTCGTCAAAGGAACCAGTACCGGGCCGCCCTACA : 750
751 : GAGGCTCGGCAAGTAGGTGGTTCCCTCTTCAGCCAGACCTCGCCATGCGAC : 800
801 : TCCCAAGACGACCAGACTCTTATTAGGTTGCTTTCGTATCCTGATTTAAA : 850
851 : AGGTAAAATAAGAGCCTGTACAACAAGCCTCCTGTCTAATAAAAATAAGA : 900
901 : GAAATAT : 907

```

**Figure 6.** Nucleotide sequence of *g-CMG1* fragment. The 5' and 3' untranslated regions are underlined. The intron sequence is indicated by dashed line.

```

1      :TCAGTGCAGAGGGAGAGCCTCGAAGTCGGTGGTCCTTGTTCCCTTCTGCTC: 50
51     :TCTGTCGAGTCCAGCGAAATGGTTGCGGTCCGATTGGTAAAGGAGGTTTA: 100
      -----
101    :AGTTCAGGCGGATTTAACTCTTGGTAAAGTGCAGAGCTTATCCTGTGCA: 150
      ----- Intron 1 -----
151    :AATATTGTTTTTTTTTCGTATAGGGGATCCGATTTTTTTTTTTTCTTTAT: 200
      ----- Intron 1 -----
201    :TAATTTTCCAATGCTTCTGACGCTGTATGCATTATTCATTTGTCTTGTGT: 250
      ----- Intron 1 -----
251    :GTCCCCTCCGAACAAAACCTGATTGATTCGAGAGTAGTAGTAAAGCGAACA: 300
      ----- Intron 1 -----
301    :TAGGAAGTTTTTCTAAACATCGAAGTCATAATCATTATATCAATACGGCC: 350
      ----- Intron 1 -----
351    :ACAAAATCCTGTAAGGGTCTCAGGACTGACCTCGGACATGAATAATTTCA: 400
      ----- Intron 1 -----
401    :GCAAGGTCCAATTTTTTTCTTAGTTACAGATGTTGATACTTCTTGGAAC: 450
      ----- Intron 1 -----
451    :AAACCGTCCTCTGAAGTTAACTCCTCTCCCTCTGCCAACAGGTGCAGTCA: 500
      ----- Intron 1 -----
501    :GCTGTTCTTGTGTCTTGTGCTGGTGGCACTTCCGGCCTGTGTACAACTTC: 550
551    :TGAAAACACGAATGAAATACCGGCGTCCATTCTTTCTTCCCCTGGGATTC: 600
601    :CCTAACAGAGACCAAAGCATAAGCAAACGTTTATATTGTTCAATTCTGGC: 650
651    :ACGGGGTCTACGACCGCGAACTTCTTGTAAGGTTGACCGCGTGTGCGAA: 700
701    :GACTGCTACAACCTCTACCGGACGTCGGAGTGGCGGCCGAATGCAGGTAT: 750
      -----
751    :GTTATTCTAATTCTTACATAAAAAATGTCAACAACCACTTTATACAGCAGC: 800
      ----- Intron 2 -----
801    :GAGTATCTGTAATCGCCTGTCAAGATCAGATGTGATAGTATGCTGGGTAA: 850
      ----- Intron 2 -----
851    :ATACTGTTATCATTTAATAGGTGATTTTTTAATCACTATAAAACCTTTA: 900
      ----- Intron 2 -----
901    :TTTAGATTCCAAGGATCTGAGTAAAATCAACTTTATTTCCCTTTTCTCTT: 950
      ----- Intron 2 -----
951    :GCATTCTATTTTCATTCTTTACCTACTTACATGGTTACATAGAACC: 1000
      ----- Intron 2 -----
1001   :TTTACATATTGCAGGAGTAACGTGTTCCACAACGAGGTGTTCTGTACTG: 1050
      -----
1051   :CGTGGACTACATGTTCCGGCCTCGCCAGAGGAACCAGTACCGGGCCGCC: 1100
1101   :TCCAGAGGCTCGGCAAGTAGGCGGTTTCGTCTTCGGTCAACCCTTCCTGCG: 1150
1151   :GGGACGCTCGCCGTGAGATTTTTTCTAGGCTCTCCTTCGGTTGCTTCTGT: 1200
1201   :ATCCTAAGTTAGAAACGGATAACATATTGACGTCGAGGTGTTTCATGTAC: 1250
1251   :AGTGTCTTCATAAAGCAG : 1268

```

**Figure 7.** Nucleotide sequence of *g-CMG2* fragment. The 5' and 3' untranslated regions are underlined. The sequences of the two introns are indicated by dashed line.

## **Research Output จากโครงการ**



## 6. Research Output จากโครงการ

### 6.1 ผลงานตีพิมพ์ในวารสารวิชาการนานาชาติ

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## 6.2 การเสนอผลงานในที่ประชุมวิชาการนานาชาติ

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7. Leelamanit, W., Chuethong, J. Boonyom, R., Panyim, S. Polonis, VR. and Nitayaphon, S. Genetic variability of subtypeE HIV-1 protease from therapy-naïve Thai patients. Joint International Tropical Medicine Meeting 1999. 4-6 August 1999, Bangkok, Thailand.
8. Leelamanit, W., Leutrakul, T., Jiratchariyakul, M., Ponglikitmongkol, M., Polonis, VR. and Panyim, S. Antitumor and anti HIV proteins extracted from *Momordica charantia* cultivated in Thailand. Joint International Tropical Medicine Meeting 1999. 4-6 August 1999, Bangkok, Thailand.

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### 6.3 การเสนอผลงานในที่ประชุมวิชาการในประเทศ

1. Senapin, S., Panyim, S. and Tungpradabkul, S. A new type of flagellin gene in *Pseudomonas putida*. The 24<sup>th</sup> Congress of Science and Technology of Thailand, 19-21 October 1998, Bangkok, Thailand.
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  8. Attsart, P., Kerbundit, S. and Miloslav, J. Determination of 3' end sequence of Thai isolated PRV RNA type W. 24<sup>th</sup> Congress on Science and Technology of Thailand, 19-21 October 1998, Bangkok, Thailand.
  9. Phucharoen, J. and Panyim, S. CyclinE and/or c-Jun overexpression affects cell growth, cell death and cell cycle progression in mouse fibroblast. 25<sup>th</sup> Congress on Science and Technology of Thailand, 20-22 October 1999, Pitsanulok, Thailand.
  10. Phucharoen, J. Butraporn, R. and Panyim, S. Production and Characterization of cyclinE transgenic mice and cyclinE transfected fibroblast : functional studies of cyclinE in vivo and in vitro. 25<sup>th</sup> Congress on Science and Technology of Thailand, 20-22 October 1999, Pitsanulok, Thailand.
  11. Chaicherdsakul, T., Boonchouy, C., Udomkit, A., Wilairat, P., Panyim, S. and Sonthyanon, B. Partial characterization of mitochondrial DNA of black tiger prawn (*Penaeus monodon*) 25<sup>th</sup> Congress on Science and Technology of Thailand, 20-22 October 1999, Pitsanulok, Thailand.
  12. Sithimonchai, S., Noonpakdee, W. and Panyim, S. Catalase-genetically-modified *L.plantarum* in NHAM. 25<sup>th</sup> Congress on Science and Technology of Thailand, 20-22 October 1999, Pitsanulok, Thailand.

#### **6.4 การประชุมวิชาการอณูชีววิทยาประจำปี**

1. “งานวิจัยด้านอณูชีววิทยา ครั้งที่ 1” 25 กันยายน 2540 ณ สถาบันอณูชีววิทยาและพันธุศาสตร์ มหาวิทยาลัยมหิดล ศาลายา มีผู้เข้าร่วมประชุม 42 คน และผู้เสนอผลงาน 10 คน
2. “งานวิจัยด้านอณูชีววิทยา ครั้งที่ 2” 25 กันยายน 2541 ณ สถาบันอณูชีววิทยาและพันธุศาสตร์ มหาวิทยาลัยมหิดล ศาลายา มีผู้เข้าร่วมประชุม 59 คน และผู้เสนอผลงาน 10 คน

3. “งานวิจัยด้านอนุชีววิทยา ครั้งที่ 3” 1 ตุลาคม 2542 ณ สถาบันอนุชีววิทยาและพันธุศาสตร์ มหาวิทยาลัยมหิดล ศาลายา มีผู้เข้าร่วมประชุม 79 คน และผู้เสนอผลงาน 9 คน

**รายชื่อที่ปรึกษาโครงการ  
และคณะผู้ร่วมวิจัย**

รายชื่อที่ปรึกษาโครงการและคณะผู้ร่วมวิจัย

ชื่อ - นามสกุล	ตำแหน่งวิชาการ		ต้นสังกัด		ตำแหน่งในโครงการ	
	เมื่อเข้าร่วมโครงการ	ปัจจุบัน	ภาควิชา	คณะ	มหาวิทยาลัย/สถาบัน	สถานภาพปัจจุบัน
1. คร. สกกล พันธุ์ยม	ศาสตราจารย์	ศาสตราจารย์	ชีวเคมี	คณะวิทยาศาสตร์	มหิดล	หัวหน้าโครงการ นักวิจัย
2. คร. ชนันท อังศุชนสมบัติ	อาจารย์	ผู้ช่วยศาสตราจารย์	สถาบันอนุชีววิทยาและพันธุศาสตร์		มหิดล	นักวิจัย
3. คร. วิภา จิงอุพรชัย	อาจารย์	รองศาสตราจารย์	สถาบันอนุชีววิทยาและพันธุศาสตร์		มหิดล	นักวิจัย
4. คร. ตูณิ เกิดบัณฑิต	อาจารย์	ผู้ช่วยศาสตราจารย์	สถาบันอนุชีววิทยาและพันธุศาสตร์		มหิดล	นักวิจัย
5. คร. ปุรชัย สันธยานนท์	อาจารย์	ผู้ช่วยศาสตราจารย์	สถาบันอนุชีววิทยาและพันธุศาสตร์		มหิดล	นักวิจัย
6. คร. อภินันท์ อุคมกิจ	อาจารย์	อาจารย์	สถาบันอนุชีววิทยาและพันธุศาสตร์		มหิดล	นักวิจัย
7. คร. จรยภรณ์ ฟูเจริญ	อาจารย์	อาจารย์	สถาบันอนุชีววิทยาและพันธุศาสตร์		มหิดล	นักวิจัย
8. คร. วิษณุ ตีลามนต์	อาจารย์	ผู้ช่วยศาสตราจารย์	ชีวเคมี	คณะเภสัชศาสตร์	มหิดล	นักวิจัย
9. คร. มรรต พงษ์ลิขิตมงคล	ผู้ช่วยศาสตราจารย์	รองศาสตราจารย์	ชีวเคมี	คณะวิทยาศาสตร์	มหิดล	นักวิจัย
10. คร. วิไล หนูนถัก	อาจารย์	ผู้ช่วยศาสตราจารย์	ชีวเคมี	คณะวิทยาศาสตร์	มหิดล	นักวิจัย
11. คร. สุนาลี ศังประคัมกุล	อาจารย์	ผู้ช่วยศาสตราจารย์	ชีวเคมี	คณะวิทยาศาสตร์	มหิดล	นักวิจัย
12. คร. ศราวุธ จิตรภักดี	อาจารย์	อาจารย์	ชีวเคมี	คณะวิทยาศาสตร์	มหิดล	นักวิจัย
13. น.ศ. บุญญรักษ์ บุญญวรรณ	ไม่มี	ไม่มี		คณะวิทยาศาสตร์	มหิดล	นักศึกษานิพนธ์
14. นาคำรณ รัตนสุตา	ไม่มี	ไม่มี		สถาบันอนุชีววิทยาและพันธุศาสตร์	มหิดล	นักศึกษานิพนธ์
15. น.ศ. เพ็ญนภา มานิตย์โชติพิสิฐ	ไม่มี	ไม่มี		สถาบันอนุชีววิทยาและพันธุศาสตร์	มหิดล	นักศึกษานิพนธ์



## รายชื่อที่ปรึกษาโครงการและคณะผู้ร่วมวิจัย

ชื่อ - นามสกุล	ตำแหน่งวิชาการ		ต้นสังกัด			ตำแหน่งในโครงการ	สถานภาพปัจจุบัน
	เมื่อเข้าร่วมโครงการ	ปัจจุบัน	ภาควิชา	คณะ	มหาวิทยาลัย/สถาบัน		
16. น.ศ. ประวีณา กิตติกุล	ไม่มี	ไม่มี	สถาบันอณูชีววิทยาและพันธุศาสตร์		มหิดล	นักศึกษาระดับปริญญาโท	อาจารย์ : จุฬาลงกรณ์มหาวิทยาลัย
17. น.ศ. กรรณกรณ์ วีระชาติ	ไม่มี	ไม่มี	สถาบันอณูชีววิทยาและพันธุศาสตร์		มหิดล	นักศึกษาระดับปริญญาโท	อาจารย์ : จุฬาลงกรณ์มหาวิทยาลัย
18. นายอรรถกฤษ์ กิตติวรการ	ไม่มี	ไม่มี	สถาบันอณูชีววิทยาและพันธุศาสตร์		มหิดล	นักศึกษาระดับปริญญาโท	อาจารย์ : มหาวิทยาลัยมหิดล
19. น.ศ. ธัญญา คงจินดา	ไม่มี	ไม่มี	สถาบันอณูชีววิทยาและพันธุศาสตร์		มหิดล	นักศึกษาระดับปริญญาโท	นักวิจัย : ศูนย์พันธุวิศวกรรมและเทคโนโลยีชีวภาพแห่งชาติ
20. น.ศ. กุลศิริ เจริญศัลป์	ไม่มี	ไม่มี	สถาบันอณูชีววิทยาและพันธุศาสตร์		มหิดล	นักศึกษาระดับปริญญาโท	นักศึกษาระดับปริญญาโท
21. น.ศ. จันทนา วงศ์ต้นติชน	ไม่มี	ไม่มี	สถาบันอณูชีววิทยาและพันธุศาสตร์		มหิดล	นักศึกษาระดับปริญญาโท	นักศึกษาระดับปริญญาโท
22. นายนพคุณ มุลลิน	ไม่มี	ไม่มี	สถาบันอณูชีววิทยาและพันธุศาสตร์		มหิดล	นักศึกษาระดับปริญญาโท	นักศึกษาระดับปริญญาโท
23. นายประพลทิศ ปิยะวิริยะกุล	ไม่มี	ไม่มี	สถาบันอณูชีววิทยาและพันธุศาสตร์		มหิดล	นักศึกษาระดับปริญญาโท	อาจารย์ : จุฬาลงกรณ์มหาวิทยาลัย
24. นายรังวิทย์ บุญโยม	ไม่มี	ไม่มี	สถาบันอณูชีววิทยาและพันธุศาสตร์		มหิดล	นักศึกษาระดับปริญญาโท	อาจารย์ : มหาวิทยาลัยนครสวรรค์
25. นายสุนทรศักดิ์ ชูลักษณ์	ไม่มี	ไม่มี	สถาบันอณูชีววิทยาและพันธุศาสตร์		มหิดล	นักศึกษาระดับปริญญาโท	อาจารย์ : มหาวิทยาลัยบูรพา
26. น.ศ. ศรีนพิพย์ เรืองงวาทติ	ไม่มี	ไม่มี	สถาบันอณูชีววิทยาและพันธุศาสตร์		มหิดล	นักศึกษาระดับปริญญาโท	ศึกษาต่อปริญญาเอก
27. ม.ล. เสาวรส สวัสดิ์วัฒน์	ไม่มี	ไม่มี	ชีวเคมี   คณะวิทยาศาสตร์		มหิดล	นักศึกษาระดับปริญญาเอก	นักวิจัย : มหาวิทยาลัยมหิดล
28. นายปิ่นนัท กาญจนภูมิ	ไม่มี	ไม่มี	สถาบันอณูชีววิทยาและพันธุศาสตร์		มหิดล	นักศึกษาระดับปริญญาเอก	นักศึกษาระดับปริญญาเอก
29. นายวันชัย อัสวาทกุล	ไม่มี	ไม่มี	สถาบันอณูชีววิทยาและพันธุศาสตร์		มหิดล	นักศึกษาระดับปริญญาเอก	นักศึกษาระดับปริญญาเอก

## รายชื่อที่ปรึกษาโครงการและคณะผู้ร่วมวิจัย

ชื่อ - นามสกุล	ตำแหน่งวิชาการ		ต้นสังกัด		ตำแหน่งในโครงการ	สถานภาพปัจจุบัน
	เมื่อเข้าร่วมโครงการ	ปัจจุบัน	ภาควิชา	คณะ		
30 น.ศ. พงโสภา อัครศาสตร์	นักวิจัย	นักวิจัย	สถาบันอณูชีววิทยาและพันธุศาสตร์	มหิดล	นักศึกษาระดับปริญญาเอก	นักศึกษาระดับปริญญาเอก
31. นางเพ็ญจันทร์ สมนายนนท์	นักวิจัย	นักวิจัย	สถาบันอณูชีววิทยาและพันธุศาสตร์	มหิดล	ผู้ช่วยวิจัย	ผู้ช่วยวิจัย
32. นางจุฑามาศ ศิริปาณี	นักวิทยาศาสตร์	นักวิทยาศาสตร์	สถาบันอณูชีววิทยาและพันธุศาสตร์	มหิดล	ผู้ช่วยวิจัย	ผู้ช่วยวิจัย
33. นายสมภพ ลีตะชีวะ	นักวิจัย	นักวิจัย	สถาบันอณูชีววิทยาและพันธุศาสตร์	มหิดล	ผู้ช่วยวิจัย	ผู้ช่วยวิจัย
34. น.ศ. ชนิกานต์ บุญช่วย	นักวิทยาศาสตร์	นักวิทยาศาสตร์	สถาบันวิจัยและพัฒนาวิทยาศาสตร์และเทคโนโลยี	มหิดล	ผู้ช่วยวิจัย	ผู้ช่วยวิจัย
35. นายอำนาจ ชะนะมา	นักวิทยาศาสตร์	นักวิทยาศาสตร์	สถาบันวิจัยและพัฒนาวิทยาศาสตร์และเทคโนโลยี	มหิดล	ผู้ช่วยวิจัย	ผู้ช่วยวิจัย