Characterization of Juvenile Hormone Biosynthetic Enzymes in the Mosquito, Aedes aegypti

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CHARACTERIZATION OF JUVENILE HORMONE BIOSYNTHETIC ENZYMES IN
THE MOSQUITO, *Aedes aegypti*

A dissertation submitted in partial fulfillment of the
requirements for the degree of
DOCTOR OF PHILOSOPHY
in
BIOLOGY
by
Pratik Nyati

2014
To: Interim Dean Michael R. Heithaus  
College of Arts and Sciences

This dissertation, written by Pratik Nyati, and entitled Characterization of Juvenile Hormone Biosynthetic Enzymes in the Mosquito, *Aedes aegypti*, having been approved in respect to style and intellectual content, is referred to you for judgment.

We have read this dissertation and recommend that it be approved.

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Date of Defense: November 5, 2014

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Dean Lakshmi N. Reddi  
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Florida International University, 2014
DEDICATION

This dissertation is dedicated to my parents, Dr. Prem Nyati and Brijlata Nyati for instilling in me the importance of education, hard work and a keen desire to run after the new learning. This doctoral thesis is also dedicated to my lovely better half Parul and younger brother Prakhar for a constant support and encouragement at each moment of my life.
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None of this would have been possible without my better half, Parul Maheshwari. Thank you dear Parul for your unconditional love, belief and sharing all the moments of happiness and sadness together.
ABSTRACT OF THE DISSERTATION

CHARACTERIZATION OF JUVENILE HORMONE BIOSYNTHETIC ENZYMES IN
THE MOSQUITO *Aedes aegypti*

by

Pratik Nyati

Florida International University, 2014

Miami, Florida

Professor Fernando G. Noriega, Major Professor

The juvenile hormones (JHs) are sesquiterpenoid compounds that play a central role in insect reproduction, development and behavior. They are synthesized and secreted by a pair of small endocrine glands, the *corpora allata* (CA), which are intimately connected to the brain. The enzymes involved in the biosynthesis of JH are attractive targets for the control of mosquito populations. This dissertation is a comprehensive functional study of five *Aedes aegypti* CA enzymes, HMG-CoA synthase (*AaHMGS*), mevalonate kinase (*AaMK*), phosphomevalonate kinase (*AaPMK*), farnesyl diphosphate synthase (*AaFPPS*) and farnesyl pyrophosphate phosphatase (*AaFPPrase*).

The enzyme *AaHMGS* catalyzes the condensation of acetoacetyl-CoA and acetyl-CoA to produce HMG-CoA. The enzyme does not require any co-factor, although its activity is enhanced by addition of Mg$^{2+}$. The enzyme *AaMK* is a class I mevalonate kinase that catalyzes the ATP-dependent phosphorylation of mevalonic acid to form mevalonate 5-phosphate. Activity of *AaMK* is inhibited by isoprenoids. The enzyme *AaPMK* catalyzes the cation-dependent reversible reaction of phosphomevalonate and ATP to form diphosphate mevalonate and ADP. The enzyme *AaFPPS* catalyzes the
condensation of isopentenyl diphosphate (IPP) and dimethylallyl pyrophosphate (DMAPP) to form geranyl diphosphate (GPP) and farnesyl pyrophosphate (FPP). The enzyme AaFPPS shows an unusual product regulation mechanism, with chain length final product of 10 or 15 C depending on the metal cofactor present. The enzymes AaFPPase-1 and AaFPPase-2 efficiently hydrolyze FPP into farnesol, although RNAi experiments demonstrate that only AaFPPase-1 is involved in the catalysis of FPP into FOL in the CA of A. aegypti. This dissertation also explored the inhibition of the activity of some of the JH biosynthesis enzymes as tools for insect control. We described the effect of N-acetyl-S-geranylgeranyl-L-cysteine as a potent inhibitor of AaFPPase-1 and AaFPPase-2. In addition, inhibitors of AaMK and AaHMGS were also investigated using purified recombinant proteins.

The present study provides an important contribution to the characterization of recombinant proteins, the analysis of enzyme kinetics and inhibition constants, as well as the understanding of the importance of these five enzymes in the control of JH biosynthesis rates.
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LIST OF ABBREVIATIONS AND ACRONYMS

AaEpox………………………………………………………………………………. *Aedes aegypti* epoxidase
AaFALD…………………………………………………………………………… *Aedes aegypti* farnesal dehydrogenase
AaFOLD…………………………………………………………………………… *Aedes aegypti* farnesol dehydrogenase
AaFPPase…………………………………………………………………………… *Aedes aegypti* farnesyl phosphatase
AaFPPS…………………………………………………………………………… *Aedes aegypti* farnesyl pyrophosphate synthase
AGGC………………………………………………………………………………N-acetyl-S-geranylgeranyl-L-cysteine
AaHMGR…………………………………………………………………………… *Aedes aegypti* 3-hydroxy-3-methylglutaryl-CoA reductase
AaHMGS…………………………………………………………………………… *Aedes aegypti* 3-hydroxy-3-methylglutaryl-CoA synthase
AaIPPI……………………………………………………………………………… *Aedes aegypti* isopentenyl pyrophosphate isomerase
AaJHAMT…………………………………………………………………………… *Aedes aegypti* juvenile hormone acid methyl transferase
AaMDD…………………………………………………………………………… *Aedes aegypti* mevalonate diphosphate decarboxylase
AaMK……………………………………………………………………………… *Aedes aegypti* mevalonate kinase
AaPMK…………………………………………………………………………… *Aedes aegypti* phospho mevalonate kinase
CA……………………………………………………………………………………..Corpora allata
DMAPP………………………………………………………………………………Dimethyl allyl pyrophosphate
DNA…………………………………………………………………………………. Deoxyribonucleic acid
DPM………………………………………………………………………………….Diphosphomevalonate
ETH………………………………………………………………………………….Ecdysis Triggering Hormone
FA………………………………………………………………………………….Farnesoic acid
FAL………………………………………………………………………………….Farnesal
FOL………………………………………………………………………………….Farnesol
FPP…………………………………………………………………Farnesyl pyrophosphate
GPP…………………………………………………………………Geranyl pyrophosphate
HAD………………………………………………………………haloalkanoic acid dehalogenase
HMG-CoA………………………………………………………3-hydroxy-3-methylglutaryl-CoA
IPP………………………………………………………………..Isopentenyl pyrophosphate
JH……………………………………………………………………..Juvenile hormone
MA…………………………………………………………………….Mevalonic acid
MF……………………………………………………………………..Methyl farnesoate
MVAP………………………………………………………………Mevalonate pathway
PM…………………………………………………………………….Phosphomevalonate
RNA………………………………………………………………….Ribonucleic acid
WHO…………………………………………………………….World Health Organization
Chapter 1: Introduction

1.1 *Aedes aegypti* as the principal vector of dengue fever, yellow fever and chikungunya

*Aedes aegypti* has been the focus of research in many areas owing to its global distribution and its involvement as a vector of dengue viruses, yellow fever and chikungunya. The viruses of these deadly diseases are passed on to humans through the bites of an infective female *A. aegypti* mosquito, which acquires the virus while feeding on the blood of an infected person (WHO, 2009). In the mosquito, the virus replicates in the midgut, invades the salivary gland and is ready for transmission to a new human host when the mosquito acquires a new blood meal (WHO, 2009).

Although *A. aegypti* is commonly known as the “yellow fever mosquito”, because of the availability of an effective vaccine, yellow fever is a less of a concern worldwide (Barrett and Higgs 2007); conversely dengue fever and chikungunya are presently a major health problem in tropical and subtropical regions of the planet (Barrett and Higgs 2007; Morens and Fauci 2014).

Dengue fever is endemic in more than 100 countries, with America, South-east Asia and the Western Pacific regions been the most seriously affected (Fig. 1) (Bhatt et al., 2013). In recent years, dengue has become a major international public health concern as its transmission has increased predominantly in urban and semi-urban areas. The last five decades has seen a 30-fold increase in the incidence of dengue fever (WHO, 2012). It is now estimated that over 2.5 billion people are now at risk from dengue. Some 50–100 million new infections are estimated to occur annually with a spread to previously unaffected areas. Every year hundreds of thousands of severe cases arise,
including 20,000 deaths (WHO 2012). There is a projection that by 2055 one third of the world’s population will be exposed to the risk of dengue fever. Among the factors responsible for this increase are global warming and increases in urbanization and resistance to insecticides (Hales et al., 2002). According to the World Health Organization the economic costs of dengue fever to human society is comparable to tuberculosis, malaria, hepatitis or bacterial meningitis (Gubler, 2002).

*Aedes aegypti* is also the vector of the chikungunya virus, which causes a disease clinically similar to dengue, with similar epidemiologic problems caused by the lack of vaccines and specific treatments (Morens and Fauci 2014). The name ‘chikungunya’ which means ‘to become distorted’ is derived from the Kimakonde language of the Makonde people (WHO 2014; Morens and Fauci 2014). The disease was given its name because a severe musculoskeletal pain caused affected persons to walk in a stooped posture. Chikungunya outbreaks have been identified in more than 40 countries in Asia, Africa, Europe, and America (Fig. 2). By July 11, 2014, the Pan American Health Organization had reported more than 355,000 suspected and confirmed cases of chikungunya fever from different jurisdictions in the Americas, with continuing local transmission and epidemic spread (Morens and Fauci 2014)

Over the last several years, A. aegypti transmitted diseases have caused periodic epidemics in tropical as well as subtropical regions of the world. Controlling the vector is the only way to reduce the burden of these deadly diseases. There are mainly four classes of insecticides that are available for public health purposes, namely, organ chlorides, organophosphates, carbamates, and pyrethroids (WHO 2013). The extensive use of vector control insecticides has contributed to the development of insecticide resistance,
cross resistance and adverse effects on the environment and human health. A better understanding of the biology of development and reproduction of mosquitoes might contribute to the selection of new targets for the development of better effective, safer and target specific insecticides.

Juvenile hormones (JHs) are lipophilic, acyclic sesquiterpenoids, synthesized by the corpora allata (CA), a pair of endocrine glands connected to the brain (Tobe et al., 1985). JH plays a central role in insect development, reproduction, diapause and polyphenisms (Goodman and Granger, 2005; Goodman and Cusson, 2012); therefore biosynthesis of JH has been considered as an attractive target for the chemical control of insects (Cusson et al., 2013). A number of recent reviews have summarized the current knowledge on JH biosynthesis in insects (Goodman and Cusson, 2012), as well as its potential as a target for insecticide discovery (Cusson et al., 2013).

1.2 Brief history on JH research and the discovery of JH structures.

Sir Vincent Brian Wigglesworth in 1934 wrote in his famous article named “The physiology of ecdysis in Rhodnius prolixus (Hemiptera). II. factors controlling molting and 'metamorphosis', this visionary statement: “The absence of metamorphosis in normal nymphs before the fifth stage must therefore be due to an inhibitory factor or hormone in the blood” (Wigglesworth, 1934). He also stated that the source of this “inhibitory hormone” was the corpora allata (CA). Afterward, Hans Piepho revealed that the formation and nature of the cuticle of the Lepidoptera Galleria was under hormonal control (Piepho, 1938). The work by Carroll Williams in the 1950s indicated that males of the Hyalophora cecropia moth contained a store of lipoidal “golden oil” with JH activity (William, 1956). William discovery initiated a series of studies trying to identify
JH homologs in different insects, since an active extract of the JH was then available that worked on most insect species (Gilbert et al., 2000).

The first JH (JH I) was identified from the lipid extract of the silkworm, *Hyalophora cecropia* (Roller et al., 1967) (Fig. 1). A second JH homologue (JH II) was also identified in *H. cecropia* extracts, differing from JH I by the presence of a methyl group at C7 (Meyer et al., 1968). A third JH homologue (JH III) was identified from the CA of the tobacco hornworm, *Manduca sexta* (Judy et al., 1973). JH III is the most ubiquitous homologue in insects, and also the JH form present in *Aedes aegypti* (Schooley and Baker, 1985). JH 0 and 4-methyl JH I (iso JH 0) were identified from eggs of *M. sexta* (Bergot et al., 1981) (Fig. 1). A JH III form with a second epoxide substitution at C6, C7 (JH III bisepoxide or JHB₃) is present in *Drosophila melanogaster* (Richard et al., 1989) (Fig. 1). Finally, a skipped bisepoxide (JHSB₃), with a second epoxide substitution at C2, C3, rather than at C6, C7 as in JHB₃, was identified from the pentatomid, *Plautia stali* (Kotaki et al., 2009) (Fig. 1). JH analogs (JHA) such as methoprene, pyriproxifen and fenoxycarb are compounds that mimic the action of JH and have been employed as control agents for mosquitoes, flies, stored-product pests, fleas and fire ants (Goodman and Cusson, 2012) (Fig. 1).

1.3 Juvenile hormone biosynthesis

1.3.1 Biosynthetic pathway

The JH biosynthetic pathway can be divided into two distinct biosynthetic units, the early steps and the late steps (Fig. 2) (Belles et al., 2005). The early steps follow the mevalonate pathway (MVAP) to form farnesyl pyrophosphate (FPP). In the MVAP, acetyl-CoA undergoes a series of enzymatic reactions to form the 5-carbon compound
isopentenyl-pyrophosphate (IPP). A sequential head to tail condensation of three 5C units is used to synthesize the 15-carbon farnesyl pyrophosphate (FPP) (Belles et al., 2005). FPP is a precursor of cholesterol in many organisms. Insects do not synthesize cholesterol de novo because they lack the genes encoding the enzymes required for the production of cholesterol from FPP, including squalene synthase and other subsequent enzymes of the sterol branch (Clark and Bloch, 1959). FPP is shunted into other pathways in insects, including synthesis of ubiquinone and dolichol, protein prenylation, and pheromone synthesis.

The late steps of JH biosynthesis (JH-branch) include the hydrolysis of FPP to farnesol (FOL) (Cao et al., 2009; Nyati et al., 2013), which is then successively oxidized to farnesal (FAL) by an alcohol dehydrogenase, and to farnesoic acid (FA) by an aldehyde dehydrogenase (Mayoral et al., 2009a; Rivera-Perez et al., 2013) (Fig. 2). The last two steps vary in different insect orders. In Lepidoptera, a C-10, 11 epoxidation by a P450 monooxygenase converts the FA to the epoxy acid (JH acid or JHA), which is then methylated by methyltransferase (JHAMT) to form the methyl ester. In Orthoptera and Dictyoptera, epoxidation follows methylation and this is also the case in mosquitoes (Defelipe et al., 2012; Noriega, 2014).

Expression of the first three enzymes of the JH-branch, namely, FPP phosphatase (FPPase), farnesol dehydrogenase (FOLD) and farnesal dehydrogenase (FALD), is not restricted to the CA (Nyati et al., 2013; Mayoral et al., 2009a; Rivera-Perez et al., 2013); but are expressed in several tissues, where they might play a significant role in cellular activities such as proliferation, apoptosis, signal transduction, and vesicular transport (Joo and Jetten, 2010; Pechlivanis and Kuhlmann, 2006). In contrast, the last two enzymes of
the JH-branch, the JHAMT and epoxidase (Epox) are likely exclusive for JH biosynthesis and therefore are highly expressed in the CA (Mayoral et al., 2009b; Nouzova et al., 2011).

1.3.2 Enzymes: Biochemical characterization

1.3.2.1 Acetoacetyl-CoA thiolase (EC 2.3.1.9)

It catalyzes the condensation of two molecules of acetyl CoA to form acetoacetyl CoA. Its activity is yet to be characterized in insects.

1.3.2.2 HMG-CoA synthase (EC 2.3.3.10)

HMG-CoA synthase (HMGS) catalyzes the condensation of acetoacetyl-CoA and acetyl-CoA into 3-hydroxy-3-methylglutaryl-coenzyme A (HMG-CoA). In insects HMGS has been biochemically characterized from the cockroach, *Blattella germanica* (Buesa et al., 1994; Casals et al., 1996) and *A. aegypti* (chapter 4). In both insects HMGS does not require a co-factor for activity; however the activity of the recombinant enzymes was enhanced by adding Mg$^{2+}$ (Buesa et al., 1994; chapter 4). Hymeglusin, a specific $\beta$-lactone inhibitor of the vertebrate HMGS (Greenspan et al., 1987), is an inhibitor of the HMGS enzyme activity in crude extracts from thorax and abdominal carcass of *A. aegypti* (chapter 4).

1.3.2.3 HMG-CoA reductase (EC 1.1.1.34)

HMG-CoA reductase (HMGR) reduces HMG-CoA to mevalonic acid. The activity of the HMGR has been characterized in two cockroach species: *Blattella germanica* (Casalet al., 1996) and *Diploptera punctata* (Feyereisen and Farnsworth, 1987). The activity of HMGR strictly depends on NADPH, and it is competitively inhibited by mevinolin (Feyereisen and Farnsworth, 1987).
1.3.2.4 Mevalonate kinase (EC 2.7.1.36)

Mevalonate Kinase (MK) catalyzes the phosphorylation of mevalonic acid into phosphomevalonate (PM). MK is a member of the “sugar kinase family” that includes enzymes such as galactokinase, homoserine kinase, mevalonate kinase, and phosphomevalonate (GHMP) kinase (Bork et al., 1993; Cheek et al., 2002). In insects, MK has been characterized only in *A. aegypti* (*AaMK*) (Nyati et al., in prep). The catalytic activity of *AaMK* increases in a dose response manner when Mg$^{2+}$ is added as a cofactor. *AaMK* can also utilize Mn$^{2+}$ and Co$^{2+}$ as cofactors, but the activity is much higher with Mg$^{2+}$ (Nyati *et al.*., in prep). The activity of *AaMK* is strongly inhibited by long chain isoprenoids such as geranyl-geranyl pyrophosphate (GGPP) (20 C units), farnesyl pyrophosphate (FPP) (15 C units) and geranyl pyrophosphate (GPP) (10 C units).

1.3.2.5 Phosphomevalonate kinase (EC 2.7.4.2)

Phosphomevalonate kinase (PMK) catalyzes the cation-dependent reversible reaction of PM and ATP to form diphosphate mevalonate (DPM) and ADP. Metazoans PMKs are not related to the GHMP kinases, but exhibit the typical fold of the nucleoside monophosphate (NMP) kinase family members (Smith and Mushegian, 2000). In insects, PMK has been characterized only in *A. aegypti* (*AaPMK*) (Chapter 4).

1.3.2.6 Mevalonate diphosphate decarboxylase (EC 4.1.1.33)

Mevalonate diphosphate decarboxylase (MDD) catalyzes ATP dependent decarboxylation of DPM to form isopentenyl pyrophosphate (IPP), releasing ADP and CO$_2$ as byproducts. Like MKs, MDDs are also classified as members of the GHMP kinase family (Bork *et al.*, 1993; Cheek *et al.*, 2002). Although MDDs have been cloned
in a variety of insects, and the crystal structure of non-insect proteins are known (Miziorko, 2011), its relation to JH biosynthesis is poorly characterized. Fluoromevalonate is a potent inhibitor of MDD in several lepidopteran species; inhibiting the metabolism of MA to JH very efficiently in Manduca sexta (Quistad et al., 1981).

1.3.2.7 Isopentenyl diphosphate isomerase (EC 5.3.3.2)

Isopentenyl diphosphate isomerase (IPPI) catalyzes the reversible conversion of IPP into dimethylallyl pyrophosphate (DMAPP). IPPI has been biochemically characterized from A. aegypti (Diaz et al., 2012) and two lepidopteran species: Choristoneura fumiferana and M. sexta (Sen et al., 2012). IPPIs belong to the Nudix hydrolase superfamily. IPPI activity is enhanced by Mg$^{2+}$ or Mn$^{2+}$ and inhibited by iodoacetamide in A. aegypti (Diaz et al., 2012), and by ammonium diphosphates in lepidoptera (Sen et al., 2012).

1.3.2.8 FPP synthase (EC 2.5.10)

FPP synthases (FPPSs) are short-chain prenyltransferases that catalyze two sequential coupling of IPP in a head to tail manner. The alkylation of IPP by DMAP to produce GPP occurs first; that reaction is followed by the alkylation of IPP by GPP to yield FPP. In the beetle Phaedon cochleariae, FPPS shows an unusual product regulation mechanism; it alters the chain length of its products depending on the cofactor present. The FPPS yields C$_{10}$-GPP in the presence of Co$^{2+}$ or Mn$^{2+}$, whereas it produces the longer C$_{15}$-FPP in the presence of Mg$^{2+}$ (Frick et al., 2013). A similar mechanism has been observed in A. aegypti (Rivera-Perez et al., in prep). Inhibitors of the FPPSs include bisphosphonates such as BPH-461, BPH-527 and BPH-678 (Cusson et al., 2013).
1.3.2.9 FPP phosphatase (EC 3.1.7.6)

FPP phosphatases (FPPases) catalyze the hydrolysis of FPP into FOL. The first insect FPPases were characterized in *Drosophila melanogaster* (Cao et al., 2009) and *A. aegypti* (Nyati et al., 2013). The FPPases belong to the NagD haloalkanoic acid dehalogenase family (HAD). *A. aegypti* FPPase (*Aa*FPPase) efficiently hydrolyzes FPP and GPP but not IPP (Nyati et al., 2013). Insect FPPases increase their catalytic activity in a dose response manner when Mg$^{2+}$ is used as a cofactor, and it are strongly inhibited by N-acetyl-S-geranylgeranyl-L-cysteine (AGGC) (Cao et al., 2009; Nyati et al., 2013).

1.3.2.10 Farnesol dehydrogenase (FOLD)

Two types of enzymes have been proposed for the oxidation of FOL into FAL: 1) a flavin/iron dependent alcohol oxidase activity was characterized from a CA homogenate of *Manduca sexta* (Sperry and Sen, 2001); and 2) a short chain alcohol dehydrogenase (SDR) was molecularly and biochemically characterized in the CA of *A. aegypti* (Mayoral et al., 2009a). The oxidation of farnesol by *M. sexta* CA extracts was weakly inhibited by 1, 10-phenanthroline (Sperry and Sen, 2001). In *A. aegypti* NADP$^+$ was absolutely required for FOLD activity, and NAD$^+$ or FAD$^+$ did not substitute (Mayoral et al., 2009a).

1.3.2.11 Farnesal dehydrogenase (FALD) (EC 1.2.1.B9)

A fatty aldehyde dehydrogenase (*AaALDH3*) that oxidizes FAL into FA was identified in CA of *A. aegypti* (Rivera-Perez et al., 2013). The *AaALDH3* is structurally and functionally a NAD$^+$-dependent class 3 ALDH with orthologues in many insect species (Rivera-Perez et al., 2013).
1.3.2.12 JH acid methyltransferase (JHAMT) (EC 2.1.1.15)

Recombinant JH acid methyltransferase (JHAMT) transfers a methyl group from S-adenosylmethionine (SAM) into FA to produce MF, as well as into JHA to generate JH III in all the insect species studied (Shinoda and Itoyama, 2003; Minakuchi et al., 2008; Niwa et al., 2008; Mayoral et al., 2009b). Homology modeling and docking simulations confirmed that JHAMT is capable to methylate both FA and JHA (Defelipe et al., 2011). JHAMT is an enzyme highly specific to insects hence, JHAMT specific inhibitors are expected to be excellent candidates for safe insect growth regulators.

1.3.2.13 MF epoxidase (Epox) (3.6.2.4)

MF Epox is generally consider as the last enzyme of the JH biosynthetic pathway, that catalyzes epoxidation of MF into JH in most of insects with the exception of Lepidoptera (Bhaskaran et al., 1986), Drosophila (Moshitzky and Applebaum, 1995) and Hemiptera (Kotaki et al., 2009). MF Epox uses NADPH as a cofactor, and is strongly inhibited by 1, 5-disubstituted imidazoles such as TH27 and KK96 (1-Isobutyl-5-[3-[[E)-3,7-dimethyl-2,6-octadienyl]oxy]phenyl]-1H-imidazole) (Helvig et al., 2004). Recombinant D. punctata EPOX (CYP15A1) shows higher affinity for MF, which it converted to JH III and cannot catalyze FA into JHA (Helvig et al., 2004). On the other hand Bombyx mori Epox exhibits at least 18-fold higher activity for FA than MF (Daimon et al., 2012). Therefore, the order of the methylation/epoxidation reactions depends upon the Epox substrate specificity (Defelipe et al., 2011). In Lepidoptera, Epox has a higher affinity for FA than JHAMT, so epoxidation precedes methylation; whereas in many other insects Epox cannot metabolize FA, so methylation precedes epoxidation.
1.4 JH functions

1.4.1 JH is one of the key hormones regulating growth and metamorphosis of insects

Two key hormones are responsible for the regulation of growth and metamorphosis in insects; JH and 20-hydroxyecdysone (20E) (Nijhout, 1994). Ecdysone is secreted by the prothoracic glands of preimaginal stages and the ovary of adult insects. Ecdysone is converted in periphery tissues to 20E (Goodman and Granger 2005; Hiruma and Kaneko 2013). Periodic pulses of 20E induce molts, and the nature of the molt depends on JH. The presence of JH ensures that the molt will produce another immature instar (nymphal in hemimetabolous and larval in holometabolous insects). The reduction in JH titers in the final nymph instars or larvae stages induces metamorphosis (Nijhout, 1994). Except for the higher Diptera, treatment of final-instar nymphs, larvae, or pupae with JH or JHA causes repetition of that stage (Goodman and Cusson, 2012). Conversely, experimental removal of JH at earlier instars leads to precocious metamorphosis. In some species, such as the silkworm, Bombyx mori (Tan et al., 2005) and the red flour beetle, Tribolium castaneum (Minakuchi et al., 2008), depletion of JH can even yield perfect miniature pupae and adults. For its ability to prevent differentiation without interfering with growth, JH was termed as a ‘status quo’ hormone (Goodman and Granger, 2005 and Dubrovsky, 2005).

1.4.2 JH assesses nutritional information to regulate reproduction in mosquitoes

In insect’s life one of the critical components is to correctly distribute nutritional reserves among survival, growth and reproduction (Boggs, 2009; Clifton and Noriega, 2011). JH is part of a transduction system that evaluates nutritional information and controls the development of ovaries in insects (Noriega, 2004). There are three major
sources of nutrients that are used by female *A. aegypti* during the three different phases of ovarian development. Reserves acquired from preimaginal stages, nectar feeding and blood meal are respectively used for the previtellogenesis (PVG), resting stage and vitellogenesis (VG) phases of ovary development (Briegel, 1990; Klowden, 1997; Noriega, 2004; Zhou *et al.*, 2004a, 2004b). In newly eclosed adult female *A. aegypti* JH synthesis and ovarian previtellogenic maturation are activated only if teneral nutritional reserves are elevated (Caroci *et al.*, 2004). Later, after previtellogenic maturation has been completed, JH mediates reproductive trade-offs in resting stage mosquitoes in response to nutrition (Clifton and Noriega, 2012). An adult female *A. aegypti* shows dynamic changes in JH biosynthesis, and regulation of the CA activity are quite different in the previtellogenesis, ovarian resting stage and vitellogenesis phases (Rivera-Perez *et al.*, 2014) (Fig. 3). Four distinct nutritional-dependent stages of CA activity have been described in female *A. aegypti*: inactive, active, modulated and suppressed CA on the basis of the rates of JH biosynthesis (Rivera-Perez *et al.*, 2014) (Fig. 3).

JH biosynthesis rate changes correspond well with the changes in transcript levels for most of the JH biosynthetic enzymes. Transcript levels are very low in early pupae (Nouzova *et al.*, 2011) and JH synthesis rates are undetectable in pupae 24 and 12 h before adult eclosion (Rivera-Perez *et al.*, 2014) (Fig. 3). Subsequently, in the last 6-8 h before adult emergence transcript levels for the biosynthetic enzymes commence to rise, the pupal CA becomes "competent" and starts to synthesize JH (Nouzova *et al.*, 2011). Although the CA of the newly eclosed adult female is fully competent; for the next 10-11 h it synthesizes relatively low levels of JH (10 fmol/h) (Rivera-Perez *et al.*, 2014) (Fig 3).

The brain of the mosquito plays a key role in sensing the nutrients and regulating the
activity of CA, since decapitation during 12 h after emergence prevents the rise in JH levels (Hernandez-Martinez et al., 2007). A sharp increase in JH synthesis is observed 12 h after adult emergence, which conveys information on teneral nutritional reserves and gives signal for the previtellogenic maturation of the ovaries. The process of “activation” of CA is very fast and short lasted; JH synthesis increases from 10 fmol/h to almost 100 fmol/h in 2 h, and decreases to less than 40 fmol/h in the next 2 h; remaining at this relatively high and constant rate until 24 h after emergence (Rivera-Perez et al., 2014) (Fig. 3). Later, during the ovarian resting stage, female mosquitoes are capable of synthesizing different rates of JH depending upon the amount of nutrients available. A. aegypti that were fed on a restricted 3% sucrose diet had reduced JH synthesis (Rivera-Perez et al., 2014) (Fig. 3). Finally at the VG phase; 24 h after blood feeding there is a suppression of JH synthesis (Li et al., 2003a; Rivera-Perez et al., 2014).

1.5 Regulation of JH biosynthesis

1.5.1 Role of the insulin/TOR signaling pathway

The insulin/TOR signaling pathways play a central role in the transduction of nutritional signals into cell growth and metabolism in almost all eukaryotic cells (Howell and Manning, 2011; Siddle, 2012). In D. melanogaster specific silencing of the insulin receptor (InR) in the CA completely suppresses HMG-CoA expression and renders a JH-deficient phenotype (Belgacem and Martin, 2007). In addition, D. melanogaster InR mutants have reduced JH synthesis (Tu et al., 2005). In Culex pipiens, the ability to enter into overwintering diapause is regulated by JH (Sim and Denlinger, 2008). Silencing the InR by RNAi or the downstream FOXO protein (fork head transcription factor) in C. pipiens leads to a diapause phenotype (Sim and Denlinger, 2008).
There are eight insulin like peptides (ILPs) reported in the genome of *A. aegypti*; three of them (ILP 1, 3 and 8) show expression specifically in brains of adult females (Riehle *et al.*, 2006). ILP3 shows the highest affinity binding for the *A. aegypti* insulin receptor (InR); and stimulates egg maturation by activating the insulin signaling pathway in the ovaries (Brown *et al.*, 2008). Insulin receptor tyrosine kinase activity and phosphatidylinositol 3-kinase mediate mosquito ILPs action (Riehle and Brown, 1999). Selective activators and inhibitors of insulin signaling cascades have strong effects on insulin-regulated physiological processes in mosquitoes (Riehle and Brown, 1999); for example, knockdown of the *A. aegypti* “phosphatase and tensin homolog” (*AaegPTEN*) affects insulin signaling (Arik *et al.*, 2009). Bovine insulin shows a strong and fast stimulation of JH synthesis by mosquito CA-CC dissected from 1 or 3 day old sugar fed females (Perez-Hedo *et al.*, 2013). Incubation of CA-CC with LY294002, an inhibitor of insulin signaling pathway resulted in a strong decrease in JH synthesis (Perez-Hedo *et al.*, 2013). JH biosynthetic transcript levels and JH synthesis were reduced by the systemic depletion of TOR by RNAi, as well as by administration of the TOR modulator rapamycin (Perez-Hedo *et al.*, 2013).

In *A. aegypti* starvation decreases JH synthesis via a decrease in insulin signaling in the CA. Starvation results in up regulation of the insulin receptor, which increases CA insulin sensitivity and might “prime” the gland to respond rapidly to increases in insulin levels after feeding resumption (Perez-Hedo *et al.*, 2014). During this response to starvation the synthetic potential of the CA remained unaffected, and the gland rapidly and efficiently responds to insulin stimulation by increasing JH synthesis to rates similar to those of CA from non-starved *A. aegypti* (Perez-Hedo *et al.*, 2014).
1.5.2 JH biosynthesis regulation by allatotropins and allatostatins

Allatotropins (ATs) are neuropeptides that stimulate CA activity, while allatostatins (AST) inhibit JH synthesis (Goodman and Cusson, 2012). The insect AT receptor belongs to a family of G-Protein-Coupled Receptors (GPCRs) orthologues of the vertebrate orexin/hypocretin receptors (Yamanaka et al., 2008; Horodyski et al., 2011; Vuerinckx et al., 2011; Nouzova et al., 2012). The Bombyx mori AT receptor (BmATr) is not expressed in the CA, but in the corpora cardiaca (CC); and it was suggested that AT inhibits the release of short Neuropeptide F, which in its turn inhibits JH synthesis. Thus AT exerts an indirect allatotropic effect by “derepression” (Yamanaka et al., 2008) which has not been tested in mosquitoes or additional insect species. Unlike BmATr, A. aegypti AT receptor (AeATr) is expressed in the CA (Nouzova et al., 2012). The pattern of changes of AeATr mRNA in the CA resembles the changes in JH biosynthesis, and it was suggested that the AeATr might play a role in the regulation of JH synthesis in mosquitoes (Nouzova et al., 2012); however its exact roles in vivo and the mechanisms of action of AT still need to be elucidated.

Insects AST can be grouped into three families, cockroach allatostatins (YXFGL-amide or type-A), cricket allatostatins (W2W9 or type-B), and Manduca allatostatins (PISCF or type-C) (Stay et al., 1994; Bendena et al., 1999; Stay and Tobe, 2007). The receptors for the three ASTs (A, B and C) also belong to the GPCR family with vertebrate orthologues. The AST-A receptors are related to the vertebrate galanin receptors (Kreienkamp et al., 2002), the AST-B receptors to the bombesin receptors (Johnson et al., 2003), and the AST-C receptors show similarity to the somatostatin/opioid receptors (Kreienkamp et al., 2006, Mayoral et al., 2010). AST-A
inhibits JH synthesis in *Diploptera punctata* CA, by reducing the availability of acetyl-CoA from glucose or amino acid metabolism, although the incorporation of acetate into the JH pathway remains unaffected (Sutherland and Feyereisen, 1996). Thus a major target of AST-A could be either the transport of citrate across the mitochondrial membrane or/and the cleavage of citrate to yield cytoplasmic acetyl-CoA (Sutherland and Feyereisen, 1996). Similar results have been described for the action of AST-C in mosquitoes. *Aedes aegypti* AST-C (*Aea*AST-C) showed no inhibitory activity in the presence of any of the intermediate precursors of JH indicating that the *Aea*AST-C target is located before the entry of acetyl-CoA in the pathway (Nouzova et al., in prep). Stimulation experiments using different sources of carbon (glucose, pyruvate, acetate and citrate) suggest that AST-C acts after pyruvate is transformed to citrate in the mitochondria. *In vitro* inhibition of the citrate mitochondrial carrier (CIC) mimicked the effect of *Aea*AST-C, and was overridden by addition of citrate or acetate. Treatment of the CA with a calcium modulator, thapsigargin (inhibitor of SERCA pumps) superseded *Aea*AST-C inhibition, suggesting an involvement of Ca$^{2+}$ in the *Aea*AST-C signaling pathway (Nouzova et al., unpublished manuscript).

### 1.5.3 Regulation of JH synthesis by 20-hydroxyecdysone and ecdysis triggering hormone

Twenty-hydroxyecdysone (20E) is a steroid hormone that controls molting, metamorphosis and oogenesis in mosquitoes (Margam et al., 2006; Attardo et al., 2005). 20E regulates JH synthesis in *Bombyx mori* (Gu and Chow 1996, Kaneko et al., 2011), possibly by means of a direct control on the expression of some of the JH biosynthetic enzymes (Hiruma and Kaneko 2013).
Ecdysis triggering hormone (ETH) is a small C-terminally amidated peptide that is released into the hemolymph to activate pre-ecdysis and ecdysis motor programs in the central nervous system (CNS) (Zitnan and Adams, 2012). Specialized endocrine cells called Inka cells synthesize ETH (Adams et al., 2006; Zitnan et al., 2007). The expression of the ETH gene is regulated by 20E (Zitnan and Adams, 2012). ETH receptors are expressed in the CA of the moths *Bombyx mori* and *Manduca sexta* (Yamanaka et al., 2008); so it has been suggested that ETH could play a role in the regulation of JH biosynthesis (Yamanaka et al., 2008). In *A. aegypti*, *in vitro* stimulation of the pupal CA with ETH resulted in an increase in JH synthesis; conversely silencing of *A. aegypti* ETH receptor in pupa resulted in reduced JH synthesis by the CA of one day old adult females (Areiza et al., 2014). There is an increase in the activity of JHAMT when *A. aegypti* CA is stimulated with ETH (Areiza et al., 2014). ETH increases JH synthesis in *A. aegypti* by mobilizing calcium from intracellular stores (Areiza et al., 2014).

**1.5.4 Flux control of JH synthesis rate**

The rate of JH biosynthesis is controlled by the rate of flux of isoprenoids in the pathway, which is the outcome of a complex interplay of changes in precursor pools, enzyme levels and external regulators (Fig. 4) (Nouzova et al., 2011). Changes in the nutritional status in female mosquitoes, as well as the manipulation of individual precursor pool concentrations (e.g. FOL, FAL and FA) affect the rate of JH biosynthesis (Fig. 4) (Nouzova et al., 2011; Rivera-Perez et al., 2014). There is a coordinated expression of JH biosynthetic enzymes in female pupae and adult mosquito. Positive correlations between JH synthesis and transcripts levels for the JH biosynthetic enzymes
suggest that a coordinated regulation in the transcription of the genes encoding JH biosynthetic enzymes is at least partially responsible for the changes of JH biosynthesis in the CA of mosquitoes (Nouzova et al., 2011). To understand how regulators modify JH synthesis, it is important to know their effect on the changes in the levels of all enzymes and precursor pool sizes.

The 13 distinct enzymatic steps of the JH synthetic pathway are arranged in an obligatory sequence. Each product represents the substrate for the next downstream enzyme. Enzymes are connected by metabolite pools that are common to them, for example FAL is the product of the FOLD and the substrate for FALD which shows that the pools and fluxes are critical variables in JH regulation. Fluxes are distributed to all the enzymes of the pathway rather than restricting to the rate limiting enzyme but the control of the flux differ widely in a pathway and the questions of its control cannot be answer by looking at one step in isolation or even each step in isolation (Kacser and Burns 1973). It has been postulated that in a pathway with multiple enzymes all the enzymes are in excess, so that individual amounts can be significantly reduced without considerable effect on the flux (Noriega, 2014). For example in D. punctate, inhibition of HMGR activity by one third resulted in less than 15% inhibition of JH III synthesis (Sutherland and Feyereisen, 1996), showing that this enzyme is in excess and has a low control coefficient on JH synthesis. Rate limiting bottlenecks have been proposed at different enzymatic steps in both the MVAP and JH-branch in the CA of different insects, for e.g. the activity of AaMK is controlled by feedback regulation of metabolites such as FPP and GPP operating in the downstream portions of the pathway (Nyati et al, unpublished manuscript). The low enzymatic activity of FPPase and FALD could limit the flux of
precursors and JH biosynthesis in the CA of blood-fed mosquitoes (Nyati et al., 2013; Rivera-Perez et al., 2012). In A. aegypti five enzymes: acetoacetyl-CoA thiolase, PMK, FPPase, FOLD and FALD presented overall low levels of expression in the CA (Nouzova et al., 2011; Rivera-Perez et al., 2012; Nyati et al., 2013). Under some conditions any of these enzymes could become rate limiting or bottleneck. In contrast recent studies suggest that there are multiple regulatory points in the pathway and they might change in different physiological stages (Rivera-Perez et al., 2014).

Addition of exogenous precursors such as FPP, FOL, FAL, FA, and MF stimulate JH synthesis in CA dissected from female mosquitoes. Stimulation of JH synthesis is independent of their spontaneous JH biosynthetic activity and rate of stimulation is significantly lower in suppressed CA with low JH biosynthetic activity (Nouzova et al., 2011). It is often puzzling to note that even with high endogenous pools of FOL, FAL and FA there is a limited JH synthesis in the newly emerged mosquitoes CA which can strongly be stimulated by exogenous supply of these precursors (Nouzova et al., 2011; Rivera-Perez et al., 2014). These results suggest differences in the channeling of endogenous and exogenous pools of JH precursors. Although the transcripts and catalytic activities of MVAP and JH branch enzymes are well coordinated with JH biosynthesis, the global fluctuations in the metabolite pools sizes are inversely related in the two pathways (Rivera-Perez et al., 2014; Nouzova et al., 2011). Analysis of the precursor pool size revealed remarkable modifications in the fluxes in the MVAP and JH-branch during first 24 h after adult eclosion; a 10-fold increase in JH synthesis rates by 12 h caused a striking depletion of all MVAP metabolite pools, with some of them replenished by 24 h (Rivera-Perez et al., 2014). In addition, there are examples of a reversal of the
flux in the MVP and its branches. Phosphatase activities converting DPM back into PM and PM back into MA, a kinase activity converting FOL back into FPP, and a reductase activity converting FAL back into FOL have been previously described in plants, animals and in the CA of mosquitoes (Thai et al., 1999; Rizzo and Craft, 2000; Rivera-Perez et al., 2013). Altogether these results show that the regulation of the JH biosynthesis pathway is very complex, and further studies will be necessary to reveal what factors restrict the flux into JH III at specific physiological conditions.

1.6 Dissertation objectives

Juvenile hormone (JH) plays important roles in the regulation of development, metamorphosis and reproduction of insects (Nijhout, 1994; Goodman and Cusson 2012; Noriega, 2014). The study of JH is very challenging because of the minute size of the CA and the “sticky” and unstable nature of JH. To understand the rate limiting steps and regulatory points on the regulation of JH synthesis we need to complete the molecular and biochemical characterization of the 13 enzymes involved in the pathway. Four CA enzymes IPP isomerase (Diaz et al., 2012), farnesol dehydrogenase (Mayoral et al., 2009a), farnesal dehydrogenase (Rivera-Perez et al., 2013) and methyl transferase (Mayoral et al., 2009b) have already been characterized in our laboratory. The goal of my dissertation research is to characterize five additional CA enzymes namely, HMG-CoA synthase (HMGS), mevalonate kinase (MK), phosphomevalonate kinase (PMK), farnesyl pyrophosphate synthase (FPPS) and farnesyl pyrophosphatase (FPPase), and to establish their role in the regulation of JH synthesis.
1.7 Dissertation organization

In the first chapter of my dissertation I introduce my research model organism, 
*Aedes aegypti*. The mosquito is an important vector of three deadly human viral diseases: dengue fever, yellow fever and chikungunya. The study of the hormonal regulation of development and reproduction, including a better characterization of the enzymes involved in JH synthesis might offer opportunities to identify new targets for alternative control strategies.

Chapter 2 describes the characterization of an *A. aegypti corpora allata* farnesyl phosphatase, the enzyme responsible for the conversion of farnesyl pyrophosphate (FPP) into farnesol in the biosynthesis of juvenile hormone.

Chapter 3 reports the characterization of an *A. aegypti corpora allata* mevalonate kinase, the enzyme responsible for the conversion of mevalonate into phosphomevalonate. This chapter describes for the first time the existence of a negative feedback on juvenile hormone synthesis through the action of isoprenoids on the mevalonate kinase activity.

Chapter 4 reports the partial characterization of the activities of recombinant HMG-CoA synthase, phosphomevalonate kinase and FPP synthase from *A. aegypti*

Chapter 5 summarizes the conclusions of my research and the future directions of studies aiming to complete the characterization of the enzymes of the JH biosynthetic pathway.
Figure 1: Current global burden of dengue. The map represents the cartogram of the annual number of infections for all ages, as a proportion of national or subnational geographical area (Bhatt et al., 2013).
Figure 2: Map of the countries and territories where chikungunya cases have been reported. Dark green represent the areas where chikungunya have been reported (as of August 26, 2014, map courtesy WHO, 2014).
Figure 3: Structure of juvenile hormones. Chemical structures of major naturally occurring JH homologues and commonly used JH agonists (from Goodman and Cusson, 2012).
Acetyl-CoA + Acetyl-CoA
↓ 1. Aceto-acetyl-CoA thiolase (Thiol)
Acetoacetyl-CoA (4C)
↓ 2. HMG-CoA synthase (HMGS)
3-HMG-CoA (3-hydroxy-3-methylglutaryl-CoA) (6C)
↓ 3. HMG-CoA reductase (HMGR)
Mevalonate (C6)
↓ 4. Mevalonate Kinase (MK)
5-phosphomevalonate
↓ 5. Phosphomevalonate kinase (PMK)
5-pyrophosphomevalonate
↓ 6. Mevalonate decarboxylase (MDD)
Isoprene intermediate (C5).
Isopentenyl-PP (5) x 3 7. IPP isomerase (IPPI)
↓ 8. Farnesyl Diphosphate synthase (FPPS)
Farnesyl diphosphate (FPP) (C15)
↓ 9. Farnesyl Pyrophosphatase (FPPase)
Farnesol (FOL) (C15)
↓ 10. Farnesol dehydrogenase (FOLD)
Farnesal (FAL) (C15)
↓ 11. Farnesal dehydrogenase (FALD)
Farnesoic acid (FA) (C15)
↓ 12. Methyltransferase (JHAMT)
Methyl Farnesoate (MF) (C16)
↓ 13. Epoxidase (Epox)
Juvenile Hormone III

Figure 4: Juvenile hormone III (JH III) biosynthetic pathway in mosquitoes. The biosynthetic pathway of JH III in the CA of mosquitoes is divided into two steps: early step (mevalonate pathways, MVP) and late steps (JH branch of the MVP) (modified from Belles et al., 2005). Precursors are shown in bold and connected by arrows. Enzymes are shown in italic and numbers before the enzymes refer to their positions in the pathway. Abbreviations for the enzymes are between brackets.
Figure 5: Rate of JH biosynthesis and ovarian development in female *Aedes aegypti*. Top panel: representative images of the progression of ovary development from emergence to 24 h after blood feeding. The inset in 96 h shows the lipid content of follicles from females fed 3% sugar (top) and 20% sugar (bottom). Colors for the panels match colors for the CA physiological phases described in the bottom panel. Bottom panel: JH biosynthesis by CA dissected from pupa, sugar-fed and blood-fed adult females. Hours represent times before (pupa) and after adult emergence (sugar-fed), or after blood feeding (BF). Colors represent the four distinct CA physiological phases identified: inactive or low activity CA (blue), active CA (black), modulated CA (green) and suppressed CA (red) (River-Perez *et al.*, 2014)
Figure 6: Flux model of juvenile hormone synthesis. This is a schematic representation of a working model for the control of the flux of precursors in the JH biosynthetic pathway. Precursor pools (Acetyl-CoA, S2, etc.) are represented by circles and connected by arrows. E: Enzymes are followed by a number that refers to the position in the pathway (from Nouzova et al., 2011).
1.8 References


67. Nyati P, Rivera-Perez C, Noriega FG Negative feedback by isoprenoids upon mevalonate kinase activity might regulate juvenile hormone synthesis in Aedes aegypti. In prep


77. Rivera-Perez C, Nyati P, Noriega FG *A corpora allata* prenyltransferase in mosquito displaying a metal ion substrate specificity. In prep.


Chapter 2: Farnesyl phosphatase, a corpora allata enzyme involved in juvenile hormone biosynthesis in Aedes aegypti

2.1 Abstract

The only recognized FPP phosphatase (FPPase) expressed in the corpora allata (CA) of an insect was recently described in Drosophila melanogaster (DmFPPase). A search for orthologs of the DmFPPase in Aedes aegypti led to the identification of 3 putative FPPase paralogs expressed in the CA of the mosquito (AaFPPases-1, -2, and -3). AaFPPase-1 and AaFPPase-2 were found to efficiently hydrolyze farnesyl diphosphate (FPP) into farnesol (FOL) and considered as the members of the NagD family of the Class IIA C2 cap-containing haloalkanoic acid dehalogenase (HAD) super family. Using a newly developed assay utilizing fluorescent tags, we demonstrate that AaFPPase activities were different in CA of sugar and blood-fed females. Injection of dsRNAs resulted in a significant reduction of AaFPPase-1 and AaFPPase-2 mRNAs, but only depletion of AaFPPase-1 caused a significant decrease on JH synthesis. These results suggest that AaFPPase-1 is involved in the catalysis of FPP into FOL in the CA of A. aegypti.

2.2 Introduction

Characterization of corpora allata (CA) enzymes has been hindered by the minute size of the endocrine gland; recently, the first description of an FPP phosphatase (FPPase) expressed in the CA of an insect was described in Drosophila melanogaster (Cao et al., 2009). This enzyme is a member of the haloalkanoic acid dehalogenase (HAD) super family that catalyzes phosphoryl transfer reactions on a remarkably diverse set of substrates and includes enzymes such as: phosphoesterases, ATPases,
phosphonatases, dehalogenases and sugar phosphomutases (Allen and Dunaway-Mariano, 2004; Arvind et al., 1998). HAD phosphatases employ an aspartate residue as a nucleophile in a magnesium-dependent phosphoaspartyl transferase reaction. The HAD superfamily is represented in the proteomes of organisms from all three super-kingdoms. The highly conserved structural core of the HAD enzymes consists of a α/β domain that adopts the topology typical of the Rossmann α/β folds housing the catalytic site and can be distinguished from all other Rossmanoid folds by two unique structural motifs: 1) an almost complete α-helical turn, named the ‘squiggle’, and 2) a β hairpin turn, termed the ‘flap’ (Lahiri et al., 2004; Allen and Dunaway-Mariano, 2009). The catalytic site is thus a composite of the four loops of the core domain and loop 5 of the cap domain. Whereas the core domain orchestrates the core chemistry, the cap domain functions in adapting that chemistry to a specific substrate (Lahiri et al., 2004).

The HAD superfamily can be divided into three generic classes on the basis of existence and location of a cap domain involved in substrate recognition. Class I possesses a small α-helical bundle cap between motifs I and II; Class II displays a cap between the second and third motifs; and Class III members present no cap domain (Lahiri et al., 2004). Members of the HAD phosphatase superfamily have four conserved amino acid signature motifs (Koonin and Tatusov, 1994; Arvind et al., 1998; Seifried et al., 2013). These 4 signature motifs are also well conserved in the FPPase described in Drosophila (DmFPPase) (Cao et al., 2009). Bioinformatics searches for orthologs of the DmFPPase in A. aegypti led to the identification of 3 putative FPPase paralogs expressed in the CA of the mosquito (AaFPPase-1, -2, and -3). Recombinant AaFPPase-1 and AaFPPase-2 were found to efficiently hydrolyze FPP into FOL. Different FPPase
activities were detected in CA extracts from adult female mosquitoes having diverse JH biosynthetic rates. Injection of dsRNAs resulted in a significant reduction of AaFPPase-1 and AaFPPase-2 mRNAs, but only reduction of AaFPPase-1 caused a significant decrease on JH biosynthesis. These results suggest that AaFPPase-1 is predominantly involved in the catalysis of FPP into FOL in the CA of A. aegypti.

2.3 Materials and Methods

2.3.1 Chemicals

Farnesyl diphosphate (FPP), geranyl diphosphate (GPP) and isopentenyl diphosphate (IPP) were purchased from Echelon Biosciences (Salt Lake City, UT). We purchase p-nitrophenyl phosphate from MP Bio medicals (Santa Ana, CA). The N-acetyl-S-geranylgeranyl-L-cysteine (AGGC) and N-acetyl-S-farnesyl-L-cysteine (AFC) were purchased from Cayman chemicals (Ann Arbor, MI). Taurolithocholic acid 3-sulfate was purchased from Sigma-Aldrich (St. Louis, MO).

2.3.2 Insects

Aedes aegypti of the Rockefeller strain were reared at 28°C and 80% relative humidity under a photoperiod of 16 h light: 8 h dark. A cotton pad soaked in 3% sucrose solution was provided to adults. Four-day-old female mosquitoes were membrane-fed porcine blood equilibrated to 37°C, and ATP was added to the blood meal to a final concentration of 1 mM immediately before use.

2.3.3 Expression of recombinant AaFPPases

The AaFPPase cDNAs were expressed in E. coli cells as described by Mayoral et al., 2009. Recombinant His-tagged proteins were purified using HiTrap chelating columns and PD-10 desalting columns (Amersham Pharmacia, Piscataway, NJ). Glycerol
was added to the enzyme solution (final concentration 50%), and samples were stored at -
20°C until used. Protein concentrations were determined using the bicinchoninic acid
protein assay reagent (BCA) (Pierce, Rockford, IL). Bovine serum albumin was used as a
standard.

2.3.4 Enzyme assays

2.3.4.1 Phosphatase assay

The catalytic activity of recombinant AaFPPases towards p-NPP was measured in
96 well plates as described by Cao et al., 2009. Phosphatase activities towards different
isoprenoid pyrophosphate substrates were determined using the Malachite Green
Phosphate Assay Kit (Bioassay Systems, Hayward, CA); enzymatic activities were
assayed using 40 µL reaction mixtures containing 100 mM MES, pH 6.0, 2 mM MgCl2,
substrate (150 µM) and 75 ng of enzyme. After 20 min of incubation at 37°C, the reaction
was terminated by the addition of the malachite green reagent (4:1 v/v), and 30 min later
the production of Pi was measured at 630 nm using a BioTek plate reader (BioTek,
Winooski, VT). Kinetic parameters were determined by non-linear curve fitting using the
GraphPad Prism software (San Diego, CA).

2.3.4.2 RP-HPLC analysis of FPPase catalytic products.

Production of FOL from FPP hydrolysis was analyzed by reverse-phase HPLC.
FPP (250 µM) was incubated with recombinant AaFPPase for 60 min in buffer (100 mM
MES, pH 6.0, 2 mM MgCl2). Reactions were terminated by adding 500 µl of acetonitrile.
Samples were centrifuged at 14,000 rpm for 5 min and the organic phase was recovered,
filtered and analyzed by reverse-phase HPLC on a Dionex Summit System (Dionex,
Sunnyvale, CA) equipped with a UVD 170U detector, 680 HPLC pump, TCC 100
column oven and Chromeleon software. The HPLC analysis was performed on an analytical column Acclaim 120 C18 (250 X 2.1 mm ID, particle size 5 µm) (Dionex), using isocratic elution from 0 to 20 min (acetonitrile-water, 1:1 v/v), followed by a linear gradient from 20 to 50 min (acetonitrile-water (50 to 95%, v/v) and another isocratic elution from 50 min (acetonitrile, 95%). Flow rate was 0.2 ml/min and column temperature was 25ºC. The eluate was monitored with UV (214 nm). Water or/glycerol were used in place of recombinant enzymes in negative controls.

2.3.4.3 Effect of inhibitors on AaFPPase activity

Recombinant AaFPPases were pre-incubated with different concentrations (0 to 40 µM) of putative inhibitors for 10 min and their activities were measured using the p-NPP assay. The following compounds were tested: N-acetyl-S-geranylgeranyl-L-cysteine (AGGC), N-acetyl-S-farnesyl-L-cysteine (AFC) and taurolithocholic acid 3-sulfate.

2.3.5 Quantitative real-time PCR (qPCR)

The RNA isolation and qPCR were performed as described by Nouzova et al., 2011. The primers and probes for the house keeping gene 60S ribosomal protein rpL32 and AaFPPase transcripts are included in Table 1.

2.3.6 RNAi experiments

Synthesis and microinjections of double-stranded RNA (dsRNA) were performed as described by Perez-Hedo et al., 2014. The AaFPPases and YFP (yellow fluorescent protein) target sequences for dsRNA synthesis were amplified by PCR using the AaFPPase-i and YFP-i primers (Table 1). The resulting amplicons were diluted 50-fold, and 1 µl was used as template in PCR reactions with primers containing T7 promoter sequences (Table 1). The products from these PCR reactions were purified using a
QIAquick PCR purification kit (QIAquick sciences, Germantown, MD), and 1–2 µg of the purified DNA templates were used to synthesize dsRNAs with a Megascript RNAi kit (Ambion, Austin, TX). dsRNAs were precipitated using ammonium acetate/ethanol, and resuspended in ddH2O to a final concentration of 3–4 µg/µl. In each knockdown experiment, newly emerged female mosquitoes were cold anesthetized and injected intrathoracically with 1.6 µg of dsRNA using a Drummond Nanoject II microinjector and a micromanipulator. The effect of dsRNA was evaluated 4 days after injection, a time selected based on the analysis of dsRNA depletion experiments.

2.3.7 FPPase activity in CA extracts

FPPase activities in mosquito CA-CC (corpora allata-corpora cardiaca complex) were measured by HPLC coupled to a fluorescent detector (HPLC-FD) monitoring the production of farnesol. Glands were dissected in buffer solution (100 mM MES pH 6.0, 2 mM MgCl2). CA-CC were homogenized for 1 min, sonicated 3 min and centrifuged at 10,000 g for 10 min at 4ºC. Supernatants were recovered and used as crude extract for activity assays as previously described (Rivera-Perez et al., 2013). The reaction products were labeled with DBD-COC1 for further quantification on HPLC-FD (Rivera-Perez et al., 2012). Controls such as boiled crude extract and reactions without enzyme were included. A standard curve was constructed for the quantification of tagged farnesol.

2.3.8 JH biosynthesis assay

The amount of JH synthesized by CA-CC complexes in vitro was quantified by high performance liquid chromatography coupled to a fluorescent detector (HPLC-FD) (Rivera-Perez et al., 2012). The assay uses derivatization of JH III with a fluorescent tag with subsequent analysis by reverse phase HPLC-FD.
2.3.9 Secondary structure and phylogenetic analysis

The secondary structure for AaFPPase-1 was predicted using the protein structure homology-modeling server Swiss v.8.05 (Schwede et al., 2003; Arnold et al., 2006) and the Human pyridoxal phosphate phosphatase (2oycA), that share a similarity of 29%, as template. A Maximum-Likelihood tree was built using MEGA software version 5.1 (Tamura et al., 2011), with a bootstrapping of 1000. Pairwise deletion method was selected for the gap/missing data.

2.3.10 Statistical analysis

Statistical analyses were performed using the GraphPad Prism Software (San Diego, CA, USA). The results are expressed as means ± S.E.M. Significant differences (P < 0.05) were determined with a one-tailed student t-test or one-way ANOVA followed by a pair-wise comparison of means (Tukey’s test).

2.4 Results

2.4.1 Identification of three A. aegypti FPPases expressed in the CA

Using the sequence of a D. melanogaster FPPase (CG15739) that converts FPP into FOL (DmFPPase) (Cao et al., 2009) we screened the A. aegypti genome (Vectorbase) (Lawson et al., 2009). Eight HAD genes displaying over 48% amino acid sequence similarity were identified (Genbank accession numbers: AAEL012292, AAEL010099, AAEL010098, AAEL007097, AAEL007094, AAEL007098, AAEL007090 and AAEL009503). By examining the temporal and tissue dependent expression of the 8 HAD genes by PCR we identified 3 HADs that were expressed in the CA of adult female mosquito at appropriate times (Genbank: AAEL010099, AAEL007090 and AAEL009503) (Fig. 7); we named them AaFPPase-1, AaFPPase-2 and
AaFPPase-3 respectively, and were further considered as putative AaFPPases that could be involved in JH biosynthesis. Amino acid sequence alignments of *A. aegypti* and *D. melanogaster* FPPases revealed a number of well conserved residues typical of the HAD phosphatases, including an aspartic acid (Asp$^{36}$) that acts as the catalytic nucleophile, a serine or threonine (Ser$^{67}$) for binding the phosphate group and two aspartic acid residues (Asp$^{253}$, Asp$^{258}$) important for binding the Mg$^{2+}$ cofactor (Cronin *et al.*, 2003; Seifried *et al.*, 2013) (Fig. 8). The *AaFPPase-1* structure obtained by homology modeling exhibited the typical HAD core and cap regions, with the catalytic site as a composite of the four conserved loops of the core region and the loop 5 of the cap region (cap 2 domain) (Fig. 8).

A phylogram was generated using FPPase orthologs found in insects and human (Fig. 9). The HAD classes IA and IIA clearly separated in two distinct clusters; the main cluster comprises members of the NagD family included in the class IIA with a C2 cap domain (motif V or loop 5) located between the second and third motif. Each of these amino acid sequences contains the conserved four loops (Motif I–IV). Most of the insects phosphatases identified presented one functional HAD domain in the N-terminal of the protein; with many displaying a second incomplete HAD domain in the C-terminus. In addition, three *D. melanogaster* phosphatases had a second functional HAD domain on the C-terminal. We also identified three *D. melanogaster* sequences with a single catalytic HAD domain in the C-terminus of the proteins. Two Human HAD phosphatases (phosphoglycolate phosphatase and pyridoxal phosphatase) were also grouped in the class IIA. Finally, as outgroup we used the bi-functional human epoxy hydrolase that belongs to the Class IA, having a C1 cap located between the motif I and II; this enzyme
possess both phosphatase and epoxy hydrolase functional domains. We identified three *A. aegypti* orthologs of the epoxy hydrolase, but they only possess the epoxy hydrolase domain.

2.4.2 All *AaFPPases* hydrolyzed p-NPP, but only *AaFPPase*-1 and -2 converted FPP into farnesol

The three putative *AaFPPases* were overexpressed in *E. coli*. Recombinant His-tagged proteins (~35 kDa) were purified and phosphatase activities were measured using para-nitrophenyl phosphate (p-NPP), a chromogenic substrate for most phosphatases, including alkaline, acid, protein tyrosine and serine/threonine phosphatases. *AaFPPase*-2 (*K_m* = 315.5 ± 46.9 µM) had higher affinity for p-NPP than *AaFPPase*-1 (*K_m* = 3959.43 ± 126.78 µM). All *AaFPPases* increased their catalytic activities in a dose-response manner when Mg^{2+} was used as a cofactor (Fig. 10) reaching their maximum activity at pH 6.0 (Fig. 10), which is consistent with previous findings in fruit flies (Cao *et al.*, 2009).

The specific activities of *AaFPPases* toward isoprenoid phosphates were measured using the malachite green assay, in which the amount of released inorganic phosphate is determined by quantifying the formation of a complex between malachite green molybdate and free orthophosphate that absorbs at 620–640 nm (Veldhoven and Mannaerts, 1987). Only *AaFPPase*-1 and *AaFPPase*-2 efficiently hydrolyzed FPP into FOL (*K_m* = ~222 µM) (Table 2). *AaFPPase*-1 (*K_m* = 184.45 ± 14.16 µM) and *AaFPPase*-2 (*K_m* = 273.98 ± 2.52 µM) also efficiently hydrolyzed GPP. Both enzymes also demonstrated a low affinity for IPP (Table 1). Both enzymes displayed higher “catalytic efficiencies” for GPP than for FPP with *K_cat/K_m* specificity constants for GPP 3–4 fold higher than those for FPP (Table 2). Conversion of FPP into FOL by *AaFPPase*-1 and
*Aa*FPPase-2 was confirmed by RP-HPLC analysis (Fig. 11). For the substrates used in the present study we found no evidence that pyrophosphate was released from *Aa*FPPases catalyzed reactions. The malachite green phosphate assay does not detect pyrophosphate, but only identifies free phosphate released in solution. In addition, when we treated the products of the *Aa*FPPases catalyzed reaction with pyrophosphatase (an enzyme which cleaves a pyrophosphate into two phosphate ions) we did not detect any significant increase in the amount of free phosphate.

Two isoprenoid-derived compounds, AGGC, AFC and a lipid sulfate were evaluated as potential inhibitors of the *Aa*FPPase catalytic activity. While AGGC was a potent inhibitor of *Aa*FPPase-1 and *Aa*FPPase-2 (Fig. 12), AFC and taurolithocholic acid 3-sulfate had little effect.

**2.4.3 The CA exhibited variable FPPase activity**

*Corpora allata* extracts were able to convert FPP into FOL, with the FPPase catalytic activity increasing more than 4 fold when 2 mM MgCl2 was added (Fig. 13A). *Aa*FPPase activities were measured in CA extracts from adult female mosquitoes having three distinct JH biosynthetic conditions: basal activity (0 h or newly emerged adult), high activity (24 h sugar-fed) and suppressed activity (24 h after blood feeding). In the presence of an excess of FPP, highly active glands produced 92 fmol of FOL/CA/h, while suppressed glands produced only 45 fmol of FOL/CA/h. The CA with basal activity from newly emerged females, that produced only 12 fmol/h of JH, had quite elevated FPPase activity (210 fmol of FOL/CA/h) (Fig. 13B).
2.4.4 Tissue- and developmental-stage-specific expression of *Aa*FPPases

Quantitative real time PCR was used to analyze the tissue- and developmental-stage-specific expression of *Aa*FPPases. All three *Aa*FPPase genes were expressed in the CA, but highest transcript levels were detected in other mosquito tissues. The highest level of *Aa*FPPase-1 mRNA was detected in midgut and Malphigian tubules, while that of *Aa*FPPase-2 mRNA in Malpighian tubules and *Aa*FPPase-3 transcripts were most abundant in brain and ovaries (Fig. 14). A developmental time course of mRNA expression in the CA showed that transcripts of *Aa*FPPase-1 and *Aa*FPPase-2 were low in late pupae, increased after emergence and peaked at day one in sugar-fed mosquitoes (Fig. 15A). The *Aa*FPPase-3 transcripts levels remained relatively constant for the same period. Transcript levels for the three *Aa*FPPase genes moderately increased after blood-feeding (Fig. 15B).

2.4.5 Reduction of *Aa*FPPase-1 by RNAi caused a significant decrease on JH biosynthesis

Since *Aa*FPPase-3 did not appear to catalyze FPP, it was not further considered to have a major role in JH biosynthesis. Therefore the effect of mRNA depletion using RNAi was only studied with *Aa*FPPase-1 and *Aa*FPPase-2. Injection of dsRNA resulted in a significant reduction of *Aa*FPPase-1 and *Aa*FPPase-2 mRNAs (~80%) (Fig. 16A). Reduction of *Aa*FPPase-1 transcripts resulted in a significant reduction in JH biosynthesis when compared with CA of females treated with dsYFP or ds*Aa*FPPase-2 (Fig. 16B).
2.5 Discussion

2.5.1 Molecular and functional characterization of AaFPPases expressed in the corpora allata of mosquitoes

In this study we identified and characterized two corpora allata mosquito NagD phosphatases that are able to convert FPP into FOL. The homology model of AaFPPase-1 exhibited the typical HAD core and cap regions [Burroughs et al., 2006; Seifried et al., 2013]. The core region is considered to be a modular phosphoryl-transfer unit with the squiggle and flap motifs providing a solvent exclusion mechanism that allows HAD enzymes to alternate between “open” and “closed” conformations. The enzyme in the “open” configuration allows the substrate to enter the active site and interact with the highly conserved catalytic residues in the four core motifs and the cap (Arvind et al., 1998; Seifried et al., 2013). Upon cap closure, some residues in the cap domain enter the active site and engage in catalysis. Once the substrate is bound, the enzyme assumes a “closed” configuration and the Mg\(^{2+}\) ion in the active site interacts with the negatively charged phosphate, preparing it for nucleophilic attack by the first conserved aspartate on motif I (Seifried et al., 2013) (Fig. 17). The AaFPPase-1 and previously described DmFPPase (GC15739) (Cao et al., 2009) are both expressed in the CA, process FPP into FOL and are part of a cluster of NagD family members that contain one functional active site (HAD domain) in the N-terminus of the core unit. Additional close related NagD sequences in other insects exhibited variability on the number and location of the HAD domains; although the effect of these changes on activity and substrate specificity remains to be studied. The study of FPPases from additional insect species could help to
improve our understanding of the basis of isoprenoid phosphate binding specificity in NagD insect proteins.

2.5.2. Expression of AaFPPases genes

Previous studies in *Bombyx mori* (Kinjoh *et al*., 2007; Ueda *et al*., 2009) and *A. aegypti* (Nouzova *et al*., 2011) suggested that the transcripts for most of the JH biosynthetic enzymes were highly enriched or exclusively expressed in the CA. The last two metabolic reactions, the methylation of FA and the epoxidation of MF, are most likely exclusive for JH biosynthesis and therefore the enzymes involved (juvenile hormone acid methyl transferase and epoxidase) should be highly expressed in the CA (Nouzova *et al*., 2011). In contrast, other enzymes in the late pathway, such as the AaFPPases described in these studies, farnesol dehydrogenases (Mayoral *et al*., 2009) and farnesal dehydrogenases (Rivera-Perez *et al*., 2013) are broadly expressed in many tissues, which is not surprising since farnesol and farnesal homeostasis are vital for cells in all insect tissues. Farnesol acts as a signaling molecule in cell proliferation and apoptosis (Roullet *et al*., 1999; Joo and Jetten, 2010; Joune *et al*., 2008). Posttranslational modifications by attachment of a farnesyl group to C-terminal cysteine of target proteins by farnesyl-transferases are essential for signal transduction and vesicular transport (Pechlivanis and Kuhlmann, 2006). Farnesal dehydrogenases play key roles in the generation of fatty alcohols and fatty acids as well as in the elimination of toxic biogenic and xenobiotic aldehydes, such as those produced by oxidative damage of glycerolipids or during protein deprenylation (Jakoby and Ziegler, 1990; Rizzo and Craft, 1991; Tschantz *et al*., 2001). The presence of more than one isozyme capable of catalyzing the hydrolysis of long chain pyrophosphates in mosquitoes suggests that selection
mechanism caused duplication and diversification of members of the NagD family and facilitated the evolution of more efficient substrate specificities, as well as a better tissue and developmental regulation; essential for the critical role that these phosphatases play in every cell.

2.5.3 *Aa*FPPase-1 and JH biosynthesis

Using an HPLC-fluorescence approach, we were able to measure the changes in the production of FOL by *Aa*FPPase from CA extracts dissected from newly emerged mosquitoes, sugar-fed and blood-fed female mosquitoes. As was shown with the recombinant proteins, the FPPase activity of the CA extracts were Mg$^{2+}$ dependent, and exhibited remarkable differences among basal, highly active and depressed glands. In sugar-fed females, we found a good concordance between *Aa*FPPase-1 and -2 mRNA expressions in the CA and JH biosynthesis (Li *et al*., 2003). Although the highest transcript levels of *Aa*FPPases were found in highly active glands, the maximum enzyme activity was found in basal active glands, suggesting that the molecular basis for JH regulation is quite unique at different times during the reproductive cycle of an adult female mosquito.

We have previously described a 1000-fold difference in the levels of mRNA expression in the CA among the JH biosynthetic enzymes (Nouzova *et al*., 2011). Four enzymes presented overall low levels of expression, acetoacetyl-CoA thiolase, phosphomevalonate kinase, farnesol dehydrogenase and farnesal dehydrogenase (Nouzova *et al*., 2011; Rivera-Perez *et al*., 2013); transcripts numbers for *Aa*FPPase-1 are also low and comparable to the levels of those 4 genes. Under some conditions any of these enzymes could become rate limiting or “bottleneck”. We have reported that the
low enzymatic activity of farnesal dehydrogenase could be a restrictive factor for JH biosynthesis in the CA of blood-fed mosquitoes (Rivera-Perez et al., 2013); a similar condition might apply to AaFPase-1, the decrease in enzymatic activity detected after blood-feeding might reduce the farnesol pool to levels that could limit the flux of precursors and JH biosynthesis.

*AaFPase*-1 and -2 efficiently hydrolyzed FPP into FOL. Therefore, we selected these 2 genes for RNAi studies. Although the RNAi mediated silencing was efficient for both enzymes, we found JH biosynthesis was significantly reduced only in *AaFPase*-1 silenced mosquitoes CA, suggesting that *AaFPase*-1 is predominantly involved in JH biosynthesis.

2.6 Conclusions

A search for orthologs of a farnesyl phosphatase described in *D. melanogaster* led to the identification of two NagD *AaFPases* that are expressed in the CA of *A. aegypti* and efficiently hydrolyzed FPP into FOL. A combination of RNAi experiments and biochemical studies using CA extracts and recombinant proteins support the hypothesis that these HAD enzymes convert FPP into FOL in the CA and might be involved in JH biosynthesis in mosquitoes.
### Primers used for RT-PCR

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<th>Reverse Primer</th>
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<td>5'CGAATGTACGAACGCTGTGG 3'</td>
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<td>AAEL010099</td>
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### Primers used for Q-RT-PCR

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<td>rpL32 Reverse</td>
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<td>rpL32 Probe</td>
<td>5’GGCTTTCATGAGATGGGACA3’</td>
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### Primers used for production of dsRNA

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Table 1: Primers used for RT-PCR, Q-RT-PCR, and production of dsRNA
Figure 7: PCR analysis of the expression of eight putative phosphatase genes in the CA of adult female *Aedes aegypti*. cDNA was made from: (1) CA dissected at the time of adult emergence, (2) one day-old sugar-fed females. From left to right: AAEL012292, AAEL010099 (*Aa*FPPase-1), AAEL010098, AAEL007097, AAEL007094, AEL007098, AAEL007090 (*Aa*FPPase-2) and AAEL009503 (*Aa*FPPase-3).
Figure 8: Homology model of the overall fold of AaFPPass-1 and amino acid sequence alignment of HAD motifs and cap domains from mosquito, fruit fly and human. (A) Homology model of the overall fold of AaFPPass-1. Core region is colored in green and cap region in light blue. Motifs are indicated by colors: motif I (red), motif II (orange), motif III (pink), motif IV (yellow), cap domain (dark blue) and squiggle (chocolate). B) Molecular surface diagram illustrating the active site pocket and the cap 2 region of AaFPPass-1. Core region is colored green and cap region in light blue. Motifs are indicated by colors: Motif I (red), motif II (orange), motif III (pink), motif IV (yellow), cap domain (dark blue) and squiggle (chocolate). The two structures were constructed by PyMOL using the Human pyridoxal phosphate phosphatase (2oycA) as template. C) Amino acid sequence alignment of HAD motifs and cap domains from mosquito (AaFPPass-1, -2 and -3), fruit fly (DmFPPass), human pyridoxal phosphatase (H. sapiens_PLP) and human epoxy hydrolase (H. sapiens_sEH). The suggested functions for the motifs are: motif I is required for nucleophilic attack, motif II is responsible for substrate binding, the motif III Lys is required for stabilizing the negative charge of the reaction intermediate together with the Ser/Thr of motif II, motif IV is needed for Mg$^{2+}$ ion binding and the cap domain is involved in substrate recognition. Bold letters indicate the conserved residues in each motif. The numbers represent the amino acid positions in the sequences. “h” denotes a hydrophobic residue and “x” any residue. Accession numbers: DmFPPass (CG15739), H. sapiens_PLP (NP_064711.1) and H. sapiens_sEH (NP_001243411.1).
Figure 9: Phylogenetic analysis of HAD superfamily sequences from insects and human. Sequences are labeled with species names and accession numbers in between brackets. The bifunctional human epoxy hydrolase (NP_001243411.1) was used as outgroup. Sequences grouped in two clades. All sequences in Clade 1 are members of the NagD family included in the class IIA of HAD proteins. Sub-clades are separated by the
localization of the HAD domain and the presence of a Ser (S) or Thr (T) in motif II. The position of the functional domain is referred as N-terminus or C-terminus. Insects with two potential HAD functional domains are shown with an asterisk. Bold labels represent the *AaFPPase-1*, *AaFPPase-2* and *AaFPPase-3*. Human sequences are represented by dotted lines in the tree. All sequences in Clade 2 are epoxy hydrolases, which are members of the class IA of HAD proteins.
Figure 10: Metal dependence and optimum pH of AaFPPase. Phosphatase activity was measured using p-NPP. A) Magnesium dose-dependent increases of activities. B) Optimum pH determinations. Three different buffers were used: Sodium acetate at pH 4.5 to 5.5, MES at pH 5.5 to 7 and Tris at pH 7 to 9. Each value represents the means ± S.E.M. of three replicate assays. Relative activity is defined as a percentage of the highest value recorded.
<table>
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<tr>
<th>Substrate</th>
<th>$K_m$ (µM ± SE)</th>
<th>$V_{max}$ (min⁻¹ mg⁻¹ ±SE)</th>
<th>$K_{cat}$ (s⁻¹)</th>
<th>$K_{cat}/K_m$ (M⁻¹s⁻¹)</th>
<th>Recombinant enzymes</th>
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<tr>
<td>FPP</td>
<td>222.36 ± 11.0</td>
<td>6.45 ± 0.76</td>
<td>3.33</td>
<td>1.5 X 10⁴</td>
<td>AaFPPase-1</td>
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<td>GPP</td>
<td>184.45 ± 14.16</td>
<td>12.71 ± 0.37</td>
<td>7.92</td>
<td>4.3 X 10⁴</td>
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<tr>
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<td>&gt;900</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
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<td>FPP</td>
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Table 2: Substrate specificity of AaFPPase-1 and AaFPPase-2
Figure 11: Chromatogram of a reverse-phase high performance liquid (HPLC) analysis showing the production of farnesol from FPP by \textit{AaFPPase-1}. A) 300\textmu M FPP was incubated with \textit{AaFPPase-1} in reaction buffer for 1hr at RT. Arrow indicates farnesol (retention time 37.5 min). B) Negative control in which 1mM FPP was incubated in reaction buffer without adding enzyme for 1hr at RT.
Figure 12: Effect of the inhibitor AGGC on AaFPPase activity. Recombinant AaFPPase-1 and -2 were pre incubated with different concentrations (0 to 40 μM) of N-acetyl-S-geranylgeranyl-L-cysteine (AGGC) for 10 min and their activities were measured using the p-NPP assay.
Figure 13: FPPase activity in CA extracts. A) Effect of Mg$^{2+}$ on FPPase activity: Extracts of CA dissected from sugar-fed females 24 h after emergence were incubated with or without 2 mM MgCl$_2$. Bars represent the means ± S.E.M. of three replicates of extracts from groups of 5 CA. Asterisks denote significant difference (unpaired t-test, ***p<0.001). B) The CA exhibited variable FPPase activity: Extracts of CA dissected from newly emerged females (0), 24 h after emergence (24SF) and 24 h after blood feeding (24BF) were incubated for 1 h in the presence of an excess of FPP. Bars represent the means ± S.E.M. of three replicates of extracts from groups of 10 CA. Different letters above the columns indicate significant differences among treatments (one way ANOVA p < 0.05, with Tukey’s test of multiple comparisons).
Figure 14: Tissue specific expression of *AaFPPases*

All tissues were dissected from 3-day-old sugar-fed females, except for testis and accessory glands dissected from 3-day-old sugar-fed males. BR: brain; CA: corpora allata; SG: salivary gland; HT: heart; TG: thoracic ganglia; VG: ventral ganglia; FB: fat body; MG: midgut; HG: hindgut; MT: Malpighian tubules; OV: ovaries; TS: testis and AG: accessory gland. Each value represents the means ± S.E.M of two independent biological replicates of 10–20 tissue samples evaluated in triplicate. *AaFPPase* mRNAs are expressed as copy number of mRNA/10,000 copies of rpL32 mRNA.
Figure 15: Developmental expression of *AaFPPases*. A) Expression on pupae and sugar-fed females: mRNA was isolated from CA of pupae 24 h (-24) and 6 h before adult eclosion, newly emerged adult female (0 h), sugar-fed females 24, 48 and 72 h after eclosion. B) Expression after blood feeding. Each data point is the means ± S.E.M. of three independent biological replicates of 20 CA evaluated in triplicate. *AaFPPase* mRNAs are expressed as copy number of mRNA/10,000 copies of rpL32 mRNA.
Figure 16: dsRNA mediated knockdown of AaFPPase-1 and -2. Newly emerged female mosquitoes were injected with dsAaFPPase-1, dsAaFPPase-2 or dsYFP; 4 days later transcript and JH levels were evaluated. A) Transcript levels are expressed as % of the YFP controls. Bars represent the means ± S.E.M. of two replicates of RNA extracted from thoraxes. B) JH synthesized in vitro: CA were dissected from females injected with dsAaFPPase-1, dsAaFPPase-2 or YFP dsRNA and incubated in vitro for 4 h. JH was evaluated by HPLC-FD. Bars represent the means ± S.E.M. of four replicates of 4 CA. Different letters above the columns indicate significant differences among treatments (one way ANOVA p < 0.05, with Tukey’s test of multiple comparisons).
Figure 17: Schematic representation of the catalytic mechanism for AaFPPases. Catalysis proceeds through an aspartylphosphate intermediate. 1) Once the FPP is bound, the Mg^{2+} ion in the active site interacts with the negatively charged phosphate, preparing it for nucleophilic attack by the first conserved aspartate on motif I. 2) As a result, an acyl phosphate intermediate is formed with the carboxyl group of this aspartate and a water molecule is deprotonated by the second aspartate of motif I; hydrolyzing the acyl phosphate intermediate and returning the enzyme to the native state. 3) The enzyme forms a new complex with FMP. 4) Catalysis of FMP occurs again through an aspartylphosphate intermediate. 5) Farnesol is released and the enzyme returns to the initial state.
2.7 References


Chapter 3: Structural and biochemical characterization of a mevalonate kinase involved in juvenile hormone pathway in *Aedes aegypti*

3.1 Abstract

Mevalonate kinase catalyzes the ATP-dependent phosphorylation of mevalonic acid to form mevalonate 5-phosphate, a key intermediate in the juvenile hormone (JH) pathway. Here we report the expression, biochemical and structural characterization of an *Aedes aegypti* mevalonate kinase (*AaMK*) enzyme expressed in the *corpora allata*. Different isoprenoids were analyzed as inhibitors of the recombinant enzyme using a traditional spectrophotometric assay and a HPLC assay. We found that *AaMK* was strongly inhibited by long chain isoprenoids pyrophosphates including the 20-carbon geranyl-geranyl pyrophosphate (GGPP), the 15-carbon farnesyl pyrophosphate (FPP) and the 10-carbon geranyl pyrophosphate (GPP), all of them in the nanomolar range. Short chain isoprenoids pyrophosphates such as the 5-carbon compounds isopentenyl pyrophosphate (IPP) and dimethylallyl pyrophosphate (DMAPP) also inhibited but only in the micro molar range. Other precursors of the JH biosynthesis pathway such as phospho mevalonate (PM), diphospho mevalonate (DPM) and farnesol (FOL) were not inhibitors of MK activity. In addition we also found a feedback inhibition of *AaMK* activity by FPP and GPP in mosquito crude extracts.

3.2 Introduction

Mevalonate kinases (MK) (EC 2.7.1.36) catalyze the synthesis of phosphomevalonate (PM) by transferring the γ-phosphoryl group from ATP to the C5 hydroxyl oxygen of mevalonic acid (MA) (PM) (Fu *et al.*, 2002). The irreversible reaction requires a divalent cation and represents a key step in the production of juvenile hormone
(JH) in insects (Noriega, 2014). Originally described in the late 1950’s for its role on cholesterol synthesis (Amdur et al., 1957), MK is also involved in the synthesis of a diverse group of non-sterol isoprenoid metabolites important in various cellular functions; such as protein prenylation, protein glycosylation and cell cycle regulation (Belles et al., 2005). In addition MK dysfunctions are responsible for human inherited diseases, such as mevalonic aciduria and hyperimmunoglobulinemia D/periodic fever syndrome (Hoffman et al., 1986).

Mevalonate kinases are found in the three domains of life (Lombard and Moreira, 2010). They are members of the “GHMP kinase family”, a group of sugar kinase that originally included galactokinases, homoserine kinases, mevalonate kinases, and phosphomevalonate kinases (Bork et al., 1993; Cheek et al., 2002). The GHMP kinase family has been now extended to include other kinases such as archaea shikimate kinases, L-threonine kinases, N-acetylgalactosamine kinases, glucuronokinases, arabinose kinases, galacturonic acid kinases, mevalonate diphosphate decarboxylases (MDD), and 4-(cytidine 5-diphospho)-2-Cmethyl- D-erythritol kinases (Cheek et al., 2002). Members of the GHMP kinase family have a number of conserved amino acid signature motifs, which are involved in the binding of ATP-Mg, as well as the substrates to be phosphorylated (Houten et al., 2000; Fu et al., 2002).

Mevalonate kinases from different organisms have a homodimeric structure in solution, which is composed of identical subunits with a molecular weight ranging from 70 to 105 KDa (Yang et al., 2002; Fu et al., 2002; Andreassi et al., 2007). Kinetic studies suggest that the enzyme catalyzes an ordered sequential reaction, with mevalonate binding first to the enzyme, and with PM as the first product released after catalysis (Beytia et al.,
1970; Fu et al., 2002). Above mechanism of reaction has been postulated for the rat MK (Rattus norvegicus MK or RnMK) (Fu et al., 2002). The structure of the RnMK shows that the C5-hydroxyl group of the MA and the γ-phosphate group of ATP are located close to Aspartic acid$_{204}$ that makes a salt bridge with Lysine$_{13}$. In the complex, penta-coordinated γ-phosphate transition state is stabilized by the magnesium ion, the side chains of Glutamic acid$_{193}$ and Lysine$_{13}$, as well as the main-chain carbonyl group of Serine$_{146}$. Aspartic acid$_{204}$ act as a base, abstracting a proton from a hydroxyl group in the MA. This converts MA into an excellent nucleophile, which then attacks the γ-phosphorus of ATP. Lysine$_{13}$ is believed to maintain the aspartate residue in the deprotonated state and decrease its $pK_a$ to facilitate the proton transfer (Fu et al., 2002). The “catalytic base mechanism” is supported by site-directed mutagenesis studies. In the Homo sapiens MK (HsMK), a mutation of the critical aspartate residue (catalytic base) decreases the activity by ten thousand fold compared with the wild type (Potter and Miziorko, 1997). Similarly, the catalytic rate is reduced by a modification of the lysine residue (Potter and Miziorko, 1997; Potter et al., 1997).

Another important feature of this enzyme is that its activity is regulated by feedback inhibition by isoprenoids, such as farnesyl pyrophosphate (FPP) and geranyl pyrophosphate (GPP). The inhibition has been described for several MKs, including Homo sapiens MK (Hinson et al., 1999), Rattus norvegicus MK, (Tanaka et al., 1990) and Sus domesticus MK (Beytia et al., 1970). A similar type of inhibition is also observed for MK of plants, including Phaseolus vulgaris, Cucumis melo and Hevea latex (Gray and Kekwick, 1972), yeast such as Saccharomyces cerevisia (Oulmouden and Karst, 1991), bacteria such as Staphylococcus aureus (Voynova et al., 2003) and archaea such as
Methanococcus jannaschii (Huang et al., 1999). Isoprenoids are potent competitive
inhibitors for the binding of ATP to MKs (Hinson et al., 1997; Potter and Miziorko, 1997).
Sensitivity to feedback inhibition is much greater in eukaryotic enzymes than bacterial or
archaeal enzymes (Fu et al., 2008; Primak et al., 2011). On the basis of their feedback
inhibition, MKs can be classified into at least three different classes: 1) class I MKs:
showing feedback inhibition by isoprenoids, but not by diphosphomevalonate (DPM), 2)
class II MKs: which do not show feedback inhibition by isoprenoids, but are strongly
inhibited by DPM, and 3) class III MKs: that do not show feedback inhibition by either
isoprenoids or DPM (Primak et al., 2011).

We have characterized a mevalonate kinase activity from the CA of Aedes aegypti
(AaMK). Similarly to other animal MKs, recombinant AaMK displays a strong feedback
inhibition by long chain isoprenoids, such as geranyl-geranyl pyrophosphate (GGPP), FPP
and GPP; with Ki values of less than 1 µM. The endogenous activity of AaMK was also
strongly inhibited by adding long chain isoprenoids to the crude extract of mosquito
thoraces (containing the CA). Homology modeling was used to build the structure of
AaMK in order to elucidate the mechanism of reaction and feedback inhibition by
isoprenoids.

3.3 Materials and methods

3.3.1 Chemicals

Geranyl-geranyl pyrophosphate (GGPP), farnesyl pyrophosphate (FPP), geranyl
pyrophosphate (GPP), isopentenyl pyrophosphate (IPP), dimethyl allyl pyrophosphate
(DMAPP) and farnesol (FOL) were purchased from Echelon Biosciences (Salt Lake City,
UT). Mevalonic acid (MA), phosphomevalonate (PM), diphosphomevalonate (DPM),
phosphoenolpyruvate (PEP) and nicotinamide adenine dinucleotide reduced (NADH) were purchased from Sigma-Aldrich (St. Louis, MO). Pyruvate kinase (PK) and lactate dehydrogenase (LDH) were purchased from LEE bioscience.

3.3.2. Insects

*Aedes aegypti* of the Rockefeller strain were reared at 28 °C and 80% relative humidity under a photoperiod of 16 h light: 8 h dark. A cotton pad soaked in 3% sucrose solution was provided to adults.

3.3.3. Secondary and tertiary structure of *AaMK*

The secondary structure was predicted online using the ExPASy web tools (Peter *et al*., 1974). The mosquito deduced amino acid sequence and those selected from other insect species were aligned using ClustalW (Thompson *et al*., 1997); the MK from *Rattus norvegicus* was included in the alignment to identify some of the conserved motifs that characterize this group of enzymes.

The three dimensional model structure for *AaMK* was predicted using the crystal structure of Human MK (Protein Data Bank ID code 2r3v.3) as a template. The identity of the model and *AaMK* was 37.75%. The model was performed using the protein structure homology modeling server Swiss v.8.05 (Schwede *et al*., 2003; Arnold *et al*., 2006)

3.3.4. Expression of recombinant *AaMK*

The *AaMK* cDNA was expressed in *E. coli* cells as described by (Nyati *et al*., 2013). Recombinant His-tagged proteins were purified using HiTrap chelating columns and PD-10 desalting columns (Amersham Pharmacia, Piscataway, NJ). Glycerol was added to the enzyme solution (final concentration 50%), and samples were stored at -20 °C until
used. Protein concentrations were determined using the bicinchoninic acid protein assay reagent (BCA) (Pierce, Rockford, IL). Bovine serum albumin was used as a standard.

3.3.5 Enzyme assays

3.3.5.1 Enzyme coupled spectrophotometric assay

The catalytic activity of the AaMK was measured using a modified spectrophotometric assay that couples ADP formation to pyruvate synthesis and reduction to lactate (Primak et al., 2011). The initial rate of disappearance of NADH serves as a measurement of the phosphorylation of MA by MK. The assays were performed in triplicate in 96-well plates (BioTek, Winooski, VT) for 10 min at 30 °C. Each 100 µl of reaction mixture contained 0.5 mM phosphoenolpyruvate, 0.01 mM DTT, 0.35 mM NADH, 10 mM MgCl₂, 2 U of LDH, and 2 U of PK in 100 mM Tris-HCl pH 7.6.

The Michaelis-Menten constant, $K_{m-MA}$ was determined at a saturating concentration of ATP (5 mM); with MA concentrations ranging from 0.005 to 2.5 mM. The reactions were initiated with the addition of 150 ng of recombinant AaMK. The $K_{m-ATP}$ was determined using saturating concentrations of MA (1.25 mM) and ATP concentrations ranging from 0.005 to 5 mM. The amount of NADH oxidized to NAD⁺ was monitored continuously at 340 nm. Absorbance changes were plotted against time to determine the rates of the MK-coupled reactions. To determine steady-state kinetic parameters, data were subjected to nonlinear regression fits to the Michaelis–Menten equation using the GraphPad Prism software (San, Diego, CA).

The AaMK inhibition studies were performed in triplicate by adding to the reaction mix precursors of JH synthesis (DPM, DMAPP, IPP, GPP and FPP), as well as GGPP at
various concentrations. To determine the $K_i$ value for GPP, FPP and GGPP, we used the multicurve fits of the GraphPad Prism software.

3.3.5.2. RP-HPLC analysis of the products of MK catalysis.

The phosphorylation of MA into PM was analyzed by reverse-phase HPLC. MA (200 µM) and ATP (250 µM) were incubated with recombinant AaMK for 60 min in the reaction buffer (100 mM Tris-HCl pH 7.5, 10 mM MgCl$_2$, 0.5 mM DTT). Reactions were terminated by adding 500 µl of acetonitrile, and then vortex 1 min. Samples were centrifuged at 14,000 rpm for 5 min and the organic phase was recovered, filtered and analyzed by reverse-phase HPLC on a Dionex Summit System (Dionex, Sunnyvale, CA) as previously described (Nyati et al., 2013). Water or/and glycerol were used in place of recombinant enzyme in negative controls.

3.3.6. MK activity in the crude extract (CE) of mosquitoes

Mevalonate kinase activities from mosquito female thoraxes were measured by monitoring the production of PM using RP-HPLC. Thoraxes from 24h old 3% sugar-fed females were dissected in Aedes saline solution and transferred to a buffer solution (100 mM Tris-HCl pH 7.5, 10 mM MgCl$_2$, 0.01 mM DTT). Thoraxes were homogenized for 1 min, sonicated 3 min and centrifuged at 10,000 g for 10 min at 4 ºC. Supernatants were recovered and used as crude extract (CE) for activity assays as previously described (Nyati et al., 2013). The protein contents of the CE were measured by BCA assay. Enzymatic assay was performed as previously described using 4 mg of protein. Controls such as boiled crude extract and reactions without enzyme were included. A standard curve was constructed for the quantification of PM.
3.3.7 Statistical analysis

Statistical analyses were performed using the GraphPad Prism Software (San Diego, CA, USA). The results are expressed as means ± S.E.M. Significant differences (P < 0.05) were determined with one-way ANOVA followed by a pair-wise comparison of means (Tukey’s test).

3.4 Results

3.4.1 Molecular and structural characterization of *A. aegypti* mevalonate kinase

A single orthologue gene to the *AaMK* EST was found in the genome of *A. aegypti* (VectorBase) (Lawson *et al*., 2009). The *AaMK* gene (Vectorbase accession Num. AAEL006435) is located on supercontig 1.205, and it is composed of three exons interrupted by two introns with lengths of 68 and 57 bp. It encodes a 397 amino acid protein with an estimated molecular weight of 43.27 kDa.

Analysis of the *AaMK* structure revealed a fold consisting of a mixture of α-helices and β-sheets, which are organized into N-terminal (include amino acids 1 to 246, 358 to 397) and C-terminal (include amino acids 247 to 357) domains arranged in a V-shape that creates a central cleft, with the active site located at the base of the cleft (Fig 18). The N-terminal domain (blue in Fig. 18) is composed of ten β sheets (β1-β9 and β12) and eight α-helices (α1- α7 and α12). The C-terminal domain (green in Fig. 18) is composed of four helices (α8-α11) and two β sheets (β10 and β11) (Fig. 18). A similar structure was previously described for other MKs (Yang *et al*., 2002; Fu *et al*., 2002; Andreassi *et al*., 2007).

Amino acid sequence alignments of MKs from seven insect species including *Aedes aegypti* (*AaMK*), *Culex quinquefasciatus* (*CqMK*), *Anopheles gambiae* (*AmMK*),
Drosophila melanogaster (DmMK), Bombyx mori (BmMK), Danaus plexippus (DpMK), Apis mellifera (AmMK) and Acyrthosiphon pisum (ApMK) (with similarities between 36% and 44%) revealed the four well conserved motifs typical of MKs (Fig. 19). The Rattus norvegicus MK, a protein with a solved crystal structure (Fu et al., 2002) and a close similarity with the sequence of AaMK (31%) was also included in the analysis.

3.4.2 Expression and purification of recombinant A. aegypti mevalonate kinase

Expression of the recombinant His-AaMK protein was induced in the E. coli Rosetta strain (DE3 cells) using the pET28 vector. AaMK was purified from the soluble fraction by immobilized metal affinity chromatography; yielding a protein with a molecular weight consistent with the estimated molecular weight of 43.27 kDa (Fig. 21). It has been previously reported that inclusion of the His tag had no effect on the kinetics properties of Homo sapiens MK (HsMK) (Hinson et al., 1997) and RnMK (Chu et al., 2007); therefore the recombinant His-AaMK was used to study its enzymatic properties without cleavage of the His tag.

3.4.3 Mevalonate kinase activity

Mevalonate kinase activity was measured by using both the enzyme-coupled assay and the HPLC method (Fig. 22). The catalytic activity of AaMK increased in a dose response manner when Mg<sup>2+</sup> was used as a cofactor. Mn<sup>2+</sup> and Co<sup>2+</sup> also enhanced MK activity to a lesser degree than Mg<sup>2+</sup> (Fig. 23). The AaMK activity was investigated at different pHs. The optimum pH was found to be 7.5 to 8.0; with the enzyme exhibiting 60-70% of its optimum activity over a rather broad pH range (7 to 8.5) (Fig. 23). To determine the nucleotide specificity of MK, we first investigated the rates at which pyruvate kinase (PK) could use nucleoside diphosphates other than ADP. The activity of PK was measured
by coupling the nucleotide diphosphate-dependent production of pyruvate and reduction to lactate catalyzed by lactate dehydrogenase (LDH). Relative rates measured for three replicate assays for ADP, GDP, UDP and CDP were respectively 100, 70, 40 and 20 percent. We utilized the enzyme coupled assay to determine catalytic rates in the presence of different triphosphates phosphoryl donors. Relative rates of MK activity when ATP, GTP, TTP and CTP were used as phosphoryl donors were 100, 15, 11 and 5 percent respectively (Fig. 23).

3.4.4. Kinetic properties of AaMK

Kinetic constants were measured for the purified recombinant AaMK using the enzyme-coupled assay. The AaMK had a $V_{\text{max}}$ of 37 µmol/min/mg. The $K_m$ (affinities) for MA and ATP were 90 ± 18 µM and 140 ± 28 µM respectively. The $K_m$ of the MK for MA was comparable to those previously described in archaea, bacteria and eukaryotes (Table 3). On the other hand, the $K_m$ for ATP varies greatly in different organisms.

3.4.5 Feedback inhibition of AaMK

AaMK activity was strongly inhibited by long chain isoprenoids, including GGPP, FPP and GPP with $K_i$ values of 0.55 ± 0.28 µM, 0.44 ± 0.2 µM and 0.93 ± 0.19 µM respectively (Fig. 24). Short chain isoprenoids, such as DMAPP and IPP inhibited only in the micromolar range, with a $K_i$ value greater than 10 µM; while 6C compounds, such as PM and DPM did not inhibit AaMK activity. Similar feedback inhibitions were described for other eukaryotic MKs (Table 4).
3.4.6 Inhibition of MK activity in crude extract (CE) of mosquitoes by long chain pyrophosphates

The activity of MK in crude extracts from thoraxes dissected from sugar-feed (SF) mosquitoes 24 h after adult eclosion was measured using the enzyme-coupled assay. Addition of 100 µM FPP decreased the kinase activity by 25% (Fig 25A). The optimal conditions for catalysis of three ATP-dependent enzymes (MK, phosphomevalonate kinase and mevalonate diphosphate decarboxylase) are similar (Rivera-Perez et al., 2014); therefore we hypothesized that in our in vitro assays the catalytic reactions continued from mevalonate (MA) to isopentenyl pyrophosphate (IPP) via phosphomevalonate (PM) and diphosphomevalonate (DPM). That prevented us from detecting changes in the pools of PM and DPM by HPLC; so we measured the changes in the pools of IPP as a proxy for MK activity (Fig. 25). In the presence of 100 µM FPP we found a significant decrease in the pool of IPP, confirming the inhibitory effect of FPP on MK activity (Fig 25B).

3.5 Discussion

3.5.1 Structural and biochemical characterization of AaMK

We have biochemically characterized an A. aegypti MK involved in the synthesis of JH in the CA. An expressed sequence tag (EST) encoding AaMK was obtained from an A. aegypti corpora-allata + corpora cardiaca library, constructed and sequenced as previously described (Noriega et al., 2006). Analysis of the AaMK sequence revealed the features of a typical GHMP kinase protein. AaMK exhibits a compact globular α/β structure similar to that described for Rattus norvegicus MK (RnMK) (Fu et al., 2002). Analysis of the structure of AaMK revealed the presence of two large insertions in the N-terminal domain. The first insertion lies between β5 and α6, and includes the disordered
residues 71-110, located between the α2 and α3 (Fig. 18). The disordered residues in the sequence of AaMK are longer than in RnMK, which include residues 73-88 (Fu et al., 2002). The next insertion lies between α5 and α6 (Fig. 18). As these two insertions lie at the surface of the molecule, they are probably not involved in the catalytic function of the enzyme. However, these insertions must play a role in the stability of the MK protein (Fu et al., 2002).

The crystal structure of Methanococcus jannaschii MK (MjMK) revealed the existence of a disulfide bridge by Cys107 and Cys281 (Yang et al., 2002). The crystal structures of RnMK (Fu et al., 2002) and Streptococcus pneumoniae MK (SpMK) (Andreassi et al., 2007) were also solved. Despite the similar overall structural folding in these three enzymes, the disulfide bridge was absent in these 2 MKs. The cysteine residues forming the disulfide bond of MjMK are in motif II and motif IV respectively (Chu et al., 2007). Sequence alignments of the seven insects MK revealed that there is no cysteine residue present in the motif II (Fig. 19); therefore insects MKs do not have a disulfide bridge. The overall polypeptide folding of MKs from insects and rat are also very similar. The catalytic base aspartate and lysine are also conserved in the structure of insects; hence it is reasonable to propose a catalytic base mechanism for insects as shown in figure 20.

We measured the activity of the recombinant AaMK (Fig. 22), and compared it with different recombinant enzymes from archaea, bacteria and eukaryotes. Although it seems that the essential cation in vivo is probably Mg$^{2+}$, our investigation of recombinant AaMK suggest that Mg$^{2+}$ can be partially replaced in vitro by other divalent cations such as Mn$^{2+}$ and Co$^{2+}$ (Fig. 23). Similarly, in the process of phosphorylation, although other
nucleotide triphosphates including GTP, CTP and TTP can partially substitute for ATP as phosphoryl donors \textit{in vitro} (Fig. 23), most likely ATP is the preferential \textit{in vivo} phosphoryl donor. The results for the analysis of cofactor requirements, phosphoryl source and optimal pH of the \textit{AaMK} were in agreement with those from other previously characterized MKs (Voynova \textit{et al.}, 2003; Hedl and Rodwell, 2003).

Our kinetic studies revealed that the \textit{AaMK} $V_{\text{max}}$ for the formation of PM was comparable to that described for other MK’s, ranging from 12 to 50 µmol min$^{-1}$ mg$^{-1}$ (Table 3). The \textit{AaMK} Michaelis-Menten constants for mevalonate ($K_{M, \text{MA}}$) and ATT ($K_{M, \text{ATP}}$) were in the range of those previously described for other MKs: $K_{M, \text{MA}}$ are in a range of 19 to 236 µM; while $K_{M, \text{ATP}}$ varied between 74 to 1180 µM (Table 3).

It has been shown that the two phosphate groups of FPP compete for the binding of the phosphoryl groups of ATP; but large contributions to the inhibitor affinity are derived from binding interactions for the farnesyl moieties (Fu \textit{et al.}, 2008). The structure of animal MKs display an extra region of 25 to 50 amino acids between motif I and motif II, which are absent in bacterial and archaeal MKs. It constitutes part of the FPP binding pocket, thereby facilitating the isoprenoid binding in animal MKs (Fu \textit{et al.}, 2008). On the basis of the structure, kinetics and inhibition profile of the insect MKs, we could classify them into the same class of other animal MKs.

\textit{AaMK} was also found to be inhibited by isoprenoids; with the pattern of this inhibition resembling well those of other eukaryotic MKs (Table 4). Inhibition of the activity of MKs by isoprenoids is competitive with respect to ATP (Fu \textit{et al.}, 2008), and the eukaryotic enzymes are 1000 fold more sensitive to the inhibition by isoprenoids than bacterial and archaeal enzymes (Table 4); even if their $K_m$ values differ by only 2-3 fold.
(Table 3). The MK activity in thoraces extracts of sugar-feed (SF) mosquitoes was significantly reduced in the presence of FPP, confirming the results obtained in the studies performed with the recombinant enzyme.

### 3.5.2 Role of mevalonate kinase on JH synthesis in mosquitoes

Mevalonate kinase is a key enzyme of the MVP and therefore for the synthesis of JH (Goodman and Cusson, 2012). *Aa*MK is expressed in most adult female mosquito tissues (Fig. 26) (Nouzova *et al.*, 2011); suggesting that *Aa*MK is involved in many metabolic pathways. In the CA, there is a good correlation between *Aa*MK mRNA expression and JH synthesis (Fig. 27) (Nouzova *et al.*, 2011; Rivera-Perez *et al.*, 2014). JH synthesis is suppressed during pupae development, and therefore *Aa*MK mRNA levels are very low in the CA of early pupae. In adult females, CA *Aa*MK transcript levels and JH synthesis reached maximum values during the first day after eclosion (Fig. 27A). In addition, changes in *Aa*MK transcripts in the CA of sugar-fed and blood-fed female mosquitoes were also in agreement with the changes in JH synthesis (Fig. 27A). In larvae, pupae and adult female *Bombyx mori* the CA also displayed a good correlation between JH biosynthesis and expression of MK (Kinjoh *et al.*, 2007). Likewise changes in *Aa*MK enzymatic activity correlated well with rates of JH biosynthesis (Fig. 27B). The highest MK enzymatic activity was found in highly active glands (12h and 24h sugar-fed females), and the lowest enzymatic activity was found in the suppressed gland of blood-fed mosquitoes (Fig. 27B).

### 3.6 Conclusions

We have completed the first functional and molecular characterization of a mevalonate kinase involved in the production of JH in the CA of insects. *Aa*MK is a class
I MK that has the typical structure and functional features of other members of the GHMP kinase family. Changes in MK mRNA levels and enzymatic activity in the CA of pupa and adult female mosquitoes corresponded well with changes in JH synthesis, suggesting that AaMK transcript and activity fluctuations are at least partially responsible for the dynamic changes of JH biosynthesis during the gonotrophic cycle of female mosquitoes. The activity of AaMK was strongly inhibited *in vitro* and *in vivo* by isoprenoids such as GPP and FPP. Further studies are necessary to determine if the inhibition of MK activity by downstream metabolites might be important to regulate JH synthesis in the CA of insects.
Figure 18: Homology model of the overall fold of *AaMK* and its amino acid sequence. (A) Three dimensional structure of *AaMK*. The N-terminal is shown in cyan, C-terminal is green, and motifs are indicated by colors; motif I (red), motif II (orange), motif III (pink) and motif IV (yellow). This structure was constructed by PyMOL using the Human MK (PDB: 2r3v.3) as template. (B) N-terminal amino acids are shown in black and C terminal with green color. Helices are numbered α1 through α13 and shown inside yellow box, β-strand are numbered β1 through β12 and shown inside red box. Motifs I-IV are shown in red underlined color. Two large insertions (i.e. residues 71-110 and residues 175-194) not involved in the catalytic function of *AaMK* are shown with blue line above them.
Figure 19: Amino acid sequence alignment of conserved motifs of MK from *A. aegypti*, compared with seven different insects and rat orthologs. Amino acid sequence alignment of MK motifs from *Aedes aegypti* (*Aa*MK), *Culex quinquefasciatus* (*Cq*MK), *Anopheles gambiae* (*Am*MK), *Drosophila melanogaster* (*Dm*MK), *Bombyx mori* (*Bm*MK), *Danaus plexippus* (*Dp*MK), *Apis mellifera* (*Am*MK), *Acyrthosiphon pisum* (*Ap*MK) and *Rattus norvegicus* (*Rn*MK). Catalytic aspartate is shown in red and lysine in blue color. The suggested functions for the motifs as according to (Houten et al., 2000) are Motif I has shown to be involved in targeting MK to peroxisomes and also in the stabilization of ATP binding and protein tertiary and quaternary structure. Motif II has an ATP binding site and also its high hydrophobicity makes it a good candidate for isoprenoid binding site. Motif III has function in the activation of the enzyme. Motif IV is involved in the stabilization of the mevalonate binding. Accession numbers: *Aa*MK (AAEL006435), *Cq*MK (EDS42994.1), *Ag*MK (EAA14782.5), *Dm*MK (AGB93455.1), *Bm*MK (NP_001093299.1), *Dp*MK (EHJ79258), *Am*MK (XP_006558673.1), *Ap*MK (XP_001942835) and *Rn*MK (NP_112325.1)
Figure 20: Scheme of catalytic mechanism of AaMK. This outline for *Aedes aegypti* MK (AaMK) reaction has been inspired from previously postulated catalytic base mechanism for the rat, *Rattus norvegicus* MK (RnMK) (Fu et al., 2002). Asp^{221} makes a salt bridge with Lys^{14}; the penta-coordinated γ-phosphate transition state is stabilized by the Mg^{2+}, Glu^{210}, Ser^{159} and Lys^{14}. Asp^{204} acts as a general base, abstracting a proton from the hydroxyl group in MA. This converts MA into an excellent nucleophile, which then attacks the γ-phosphorus of ATP. Lys^{13} is believed to maintain the aspartate residue in the deprotonated state and lower its pKₐ to facilitate the proton transfer.
Figure 21: SDS PAGE (A) and Western blot (B) analysis of the recombinant mosquito AaMK. Lane contents of the gel were: 1, molecular weight standard; 2 and 3, AaMK. Molecular weight of protein standards are depicted on the Y axis.
Figure 22: Chromatogram of a reverse phase-HPLC analysis showing the production of PM from MA by AaMK. (A) Mevalonate kinase reaction in which 100 µM MA and 100 µM ATP were incubated with AaMK in reaction buffer for 1 h at 30 °C. Arrow indicates PM. (B) Negative control in which 100 µM MA and 100 µM ATP were incubated in reaction buffer without adding enzyme for 1 h at 30 °C.
Figure 23: \textit{AaMK} pH curve, effect of metal ions and specificity of the phosphoryl donor. Kinase activity was measured by enzyme coupled spectrophotometric assay; ATP and MA were held constant and data were normalized to the maximum observed reaction velocities. To ensure MK was the rate-limiting enzyme, when necessary results were verified with the following conditions: doubling the amount of MK added doubled the observed rate, doubling the amount of PK and LDH did not affect the observed rate, and doubling the PEP concentration did not affect the observed rate. (A) pH curve. Two different buffers were used: MES at pH 5.5 to 7 and Tris-HCl at pH 7 to 9. (B) Effect of metal ions. Different metal cofactors Mg\textsuperscript{2+}, Mn\textsuperscript{2+} and Co\textsuperscript{2+} were assayed for the MK activity. (C) Specificity of the phosphoryl donor. Each value represents the means ± S.E. of three replicate assays. Relative activity is defined as a percentage of the highest value recorded. 
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Figure 24: Inhibition of *Aa*MK activity by GPP (A), FPP (B) and GGPP (C). The rate of MK activity was measured at different ATP concentrations, without inhibitor and with several fixed concentration of inhibitor (indicated in bracket). Secondary plots of slope versus inhibitor concentration indicated that the $K_i$ values for GPP, FPP and GGPP were respectively 0.93 ± 0.19 µM, 0.44 ± 0.2 µM and 0.55 ± 0.28 µM.
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Table 4: Feedback Inhibition of MK using short (DPM, 6C) and long chain isoprenoids (GPP, 10 C; FPP, 15C). MjMK: Methanococcus jannaschii MK, MmMK: Methanosarcina mazei, SpMK: Streptococcus pneumonia MK, SaMK: Staphylococcus aureus MK, ScMK: Saccharomyces cerevisiae, AaMK: Aedes aegypti MK, RnMK: Rattus norvegicus MK, SdMK: Sus domesticus MK, HsMK: Homo sapiens MK. NI: No Inhibition ND: Not Determined
Figure 25: Inhibition of mevalonate pathway and *AaMK* activity by long chain pyrophosphates. (A) Inhibition of the mevalonate pathway kinase activity by FPP: 5 µg protein from thorax extracts dissected from sugar-fed females 24 h after emergence were incubated in presence and absence of 100 µM FPP in Tris-HCl buffer, pH 7.5 containing 5 mM MgCl₂ and 1 mM DTT. Kinase activities were measured by enzyme coupled assay. (B) Inhibition of the AaMK activity by FPP: 5 µg protein from thorax extracts dissected from sugar-fed females 24 h after emergence were incubated in absence and presence of 100 µM FPP in Tris-HCl buffer, pH 7.5 containing 5 mM MgCl₂, 1 mM DTT, 500 µM MA & 500 µM ATP. CE represents the endogenous levels of IPP. Bars represent the means ± SD of three replicates of extracts from groups of 3 thoraxes. Different letters above the columns indicate significant differences among treatments (one way ANOVA *p* < 0.05, with Tukey’s test of multiple comparisons).
Fig. 26: Tissue specific expression of *AaMK*. All tissues were dissected from 3-day-old sugar-fed females. BR: brain; CA: corpora allata; BR: brain; FB: fat body; MG: midgut; MT: malpighian tubules; OV: ovaries. Each value represents the means ± S.E.M of two independent biological replicates of 10–20 tissue samples evaluated in triplicate. *AaFPPase* mRNAs are expressed as copy number of mRNA/10,000 copies of rpL32 mRNA (Nouzova *et al.*, 2011).
Fig. 27: Transcript levels (A) and enzyme activity (B) of the AaMK in the developmental stage of female A. aegypti. Each data point is the means ± S.E.M. of three independent biological replicates of 20 CA evaluated in triplicate. AaMK mRNAs are expressed as copy number of mRNA/10,000 copies of rpl32 mRNA (Modified from Nouzova et al., 2011 & Rivera-Perez et al., 2014).
3.7 References


Chapter 4: Characterization of additional *corpora allata* enzymes involved in juvenile hormone biosynthesis pathway

4.1 Abstract

The present study describes the partial characterization of three additional CA enzymes involved in JH synthesis: HMG-CoA synthase (HMGS), phosphomevalonate kinase (PMK) and FPP synthase (FPPS). *Aedes aegypti* cDNAs encoding HMGS, PMK and FPPS were expressed as fusion proteins in *Escherichia coli* DH5α cells, and purified by affinity chromatography. HMGS catalyzes the condensation of acetoacetyl-CoA and acetyl-CoA to produce HMG-CoA. The reaction did not require any co-factor, although the activity of the recombinant enzyme was enhanced by adding Mg\(^{2+}\). Hymeglusin, a specific β-lactone inhibitor of the vertebrate HMGS, inhibited HMGS enzyme activity in crude extracts from thoraces and abdominal carcasses of *A. aegypti*. *Aa*PMK, a member of the nucleoside monophosphate family, catalyzes the cation-dependent reversible reaction of phosphomevalonate and ATP to form diphosphate mevalonate and ADP. Kinetics for both forward and backward reactions were determined, and we observed that the activity of *Aa*PMK was not inhibited by any of the downstream metabolites. FPPS catalyzes the synthesis of different chain length isoprenyl diphosphates depending upon availability of metal cofactors. *Aa*FPPS yielded 84% C\(_{10}\)-geranyl pyrophosphate (GPP) and 16% C\(_{15}\)-farnesyl pyrophosphate (FPP) in the presence of Co\(^{2+}\) as a cofactor, whereas it yielded 33% C\(_{10}\) GPP and 67% C\(_{15}\) FPP in the presence of Mg\(^{2+}\).
4.2 Introduction

HMG-CoA synthase

HMG-CoA synthase (HMGS) is the second enzyme of the JH pathway. It catalyzes the condensation of acetoacetyl-CoA and acetyl-CoA into HMG-CoA, releasing free CoA. Kinetic studies have established that this enzymatic step proceeds via a Bi-Bi Ping-Pong mechanism; where HMG-CoA is formed in three consecutive steps that include acetylation, condensation and hydrolysis (Fig. 28) (Miziorko and Lane, 1977; Pojer et al., 2006). Step 1 is the acetylation of acetyl-CoA to an acetyl-S-enzyme intermediate. Step 2 involves the condensation of acetoacetyl-CoA and acetyl-S-enzyme to form CoA-S-HMG-S-enzyme. Step 3 is the hydrolysis of CoA-S-HMG-S-enzyme to produce HMG-CoA (Fig. 28) (Miziorko and Lane, 1977; Pojer et al., 2006). Acetylation is the rate limiting step in the synthesis of HMG-CoA, as the rate of acetylation of HMGS by acetyl-CoA is much slower than the hydrolysis of CoA-S-HMG-S-enzyme (Miziorko and Lane, 1977).

Two isoforms of the enzyme have been identified in mammals; HMGS1 and HMGS2. HMGS1 is a cytosolic protein involved in isoprenoid/cholesterol biosynthesis, while HMGS2 is a mitochondrial protein involved in the biosynthesis of ketone bodies (Goldstein and Brown 1990; Dooley et al., 1998; Willamson et al., 1968; Dashti and Ontko, 1979). HMG-CoA reductase and HMGS1 are considered key regulatory enzymes in the biosynthesis of cholesterol (Goldstein and Brown 1990; Dooley et al., 1998). In insects, HMGS does not have any recognizable N-terminal targeting sequence to mitochondria, which suggests it is present in the cytosol (Buesa et al., 1994; Tittiger et al., 2000). Most insects have one HMGS gene, but Blattella germanica is an exception with 2 HMGS genes (HMGS1A and HMGS1B), showing 78% similarity (Buesa et al.,
1994; Belles et al., 2003). HMGS1A is intronless, and expression studies and phylogenetic analysis suggest it represent a functional retrogene derived from HMGS1B by retrotransposition (Buesa et al., 1994; Cabano et al., 1997; Casals et al., 2001). In insects HMGS has been biochemically characterized only from the cockroach Blattella germanica (Buesa et al., 1994; Casals et al., 1996).

**Phosphomevalonate kinase**

Phosphomevalonate kinase (PMK) is the second ATP-dependent enzyme in the mevalonate pathway. It catalyzes the phosphorylation of phosphomevalonate (PM) to diphosphomevalonate (DPM) (Fig. 29). PMKs are related to various human diseases, such as Zellweger syndrome and rhizomelic chondrodysplasia punctata (Braverman et al., 1997; Wanders and Romeijn, 1998). However, in mammals, PMKs have not been as well characterized as are other enzymes involved in isoprenoid biosynthesis. Animal PMKs are encoded by genes that are non-orthologous to plant, fungal, and bacterial PMKs genes (Smit and Mushegian, 2000). Analysis of the crystal structure of *Streptococcus pneumoniae* PMK confirmed that this protein belongs to the GHMP kinase family (Romanowski et al., 2002; Andreassi et al., 2009). Mammalian PMKs have been purified and characterized from a variety of tissues (Hellig and Popjak 1961a, 1961b; Bazaes et al., 1980a, 1980b; Lee and O'Sullivan 1985; Chambliss et al., 1996; Herdendorf and Miziorko, 2006, 2007). Kinetic and biophysical studies suggest that the animal PMKs belong to the nucleoside monophosphate (NMP) family, which is also known as P-loop kinases family (Herdendorf and Miziorko, 2006, 2007).

In the P-loop kinases, the β sheet is five stranded, with a highly conserved order of 23145 (Walker et al., 1982). There are two highly conserved motifs known as Walker
A (GXXXXGK/T/S) and Walker B (ZZZZD, where Z is any hydrophobic residue) (Cheek et al., 2002). The mechanism of reaction involves the Walker A motif forming a phosphate-binding loop (P-loop) that is located at the end of the first $\beta$-strand, and includes the first half-turn of the following $\alpha$-helix. The conserved lysine residue of the Walker A motif binds to and orients oxygen atoms of the $\beta$ and $\gamma$-phosphate groups of ATP (Cheek et al., 2002). The essential magnesium cation is coordinated directly by the hydroxyl group of the conserved threonine/serine of the Walker A motif, and indirectly by the conserved aspartate residue of the Walker B motif (Cheek et al., 2002).

**FPP synthase**

FPPS are prenyltransferases, also known as isoprenyl diphosphate synthases (IDS). Prenyltransferases catalyze the consecutive condensation of isopentenyl diphosphate (IPP) with allylic prenyldiphosphates, to yield products with chain lengths varying from $C_{10}$ up to many C. Based on the geometry of newly formed double bond (E or Z), and the size of the isoprenoid chain in the final product, prenyltransferases can be classified as short chain and long chain prenyltransferases. GPP synthase (GPPS) produces GPP ($C_{10}$), FPPS produces FPP ($C_{15}$) and geranyl-geranyl pyrophosphate synthase (GPPS) produces GGPP ($C_{20}$). They are classified as short chain E-prenyltransferases (Ogura and Koyama, 1998). The enzyme IDS, which produce compounds involved in respiratory quinone biosynthesis, with chain length varying from $C_{30}$ to $C_{50}$, are considered as long chain E-prenyltransferases (Okada et al., 1996). The Z-polyisoprenyl diphosphate synthases are involved in the synthesis of long chain dolichols and several other very long chain isoprenoids (Sato et al., 1999). These enzymes could be encoded by different genes, but they share the mechanism of reaction and well-defined
highly conserved motifs characteristic of proteins evolved from a common ancestor (Chen et al., 1994; Fujihashi et al., 2001).

The crystal structure of FPPS has not been determined in insects, but X-ray structures have been resolved for several organisms, including avian FPPS (Tarshis et al., 1994, 1996), Staphylococcus aureus FPPS, Escherichia coli FPPS (Hosfield et al., 2004), Trypanosoma cruzi FPPS (Gabelli et al., 2006), and human FPPS (Rondeau et al., 2006). These studies described the enzymes as homodimers, with each subunit folded in a single domain, whose central feature is a core composed of 10 α-helix surrounding a large deep cleft which is identified as the substrate binding pocket.

The reaction mechanism for FPPS is postulated as a dissociative electrophilic alkylation, which is divided into three steps as shown in figure 30. In step 1, the C1–oxygen bond in DMAPP or GPP ruptures to generate a resonance stabilized allylic cation. In step 2, this allylic cation alkylates the double bond in IPP to produce a tertiary carbocation. During step 3, hydrogen is subsequently eliminated from C2 of the IPP unit to produce a new allylic diphosphate which is one isoprene unit longer than the substrate (Fig. 30) (Poulter and Rilling, 1978; Poulter, 2006). The reaction depends upon the metal cofactor usually Mg$^{2+}$, Mn$^{2+}$ or Co$^{2+}$ (Aaron and Christianson, 2010). In the beetle Phaedon cochleariae, FPPS shows an unusual product regulation mechanism; it alters the chain length of its products depending on the cofactor present. The FPPS yields C$_{10}$-GPP in the presence of Co$^{2+}$ or Mn$^{2+}$, whereas it produces the longer C$_{15}$-FPP in the presence of Mg$^{2+}$ (Frick et al., 2013).
4.3. Methods

4.3.1. Insects

*A. aegypti* of the Rockefeller strain were reared at 28 °C and 80% relative humidity under a photoperiod of 16 h light: 8 h dark. A cotton pad soaked in 3% or 20% sucrose solution was provided to adults.

4.3.2. Tertiary structure of proteins

The three dimensional model structures for *Aa*PMK and *Aa*FPPS were predicted using the crystal structures of human PMK (Protein Data Bank ID code 3ch4.1) and avian FPPS (Protein Data Bank ID code 1ubx.1) respectively as templates. The identity of the human PMK and *Aa*PMK was 49.08 % and that of avian FPPS and *Aa*FPPS was 47.35%. The models were performed using the protein structure homology modeling server Swiss v.8.05 (Schwede et al., 2003; Arnold et al., 2006)

4.3.3. Expression of recombinant proteins

The cDNAs of *Aa*HMGS, *Aa*PMK and *Aa*FPPS were expressed in *E. coli* cells as described by Nyati *et al.*, 2013. Recombinant His-tagged proteins were purified using HiTrap chelating columns and PD-10 desalting columns (Amersham Pharmacia, Piscataway, NJ). Glycerol was added to the enzyme solution (final concentration 50%), and samples were stored at -20 °C until used. Protein concentrations were determined using the bicinchoninic acid protein assay reagent (BCA) (Pierce, Rockford, IL). Bovine serum albumin was used as a standard.
4.3.4 Enzymatic assays

4.3.4.1 PMK assays

The catalytic activity of the AaPMK was measured using a modified spectrophotometric assay that couples ADP formation to pyruvate synthesis and reduction to lactate as described in chapter 4.

4.3.4.2 FPPS assays

AaFPPS assays were carried out in a final volume of 100 μL containing 100 mM Tris-HCl (pH 7.5), 100 μM IPP, 100 μM DMAPP, and 5 mM Mg²⁺ or 1 mM Co²⁺. Reactions were started with the addition of 4 μg recombinant enzyme. Formation of FPP and GPP were monitored by transforming them into FOL and geraniol using 50 μL 3N HCL and quantifying them by reverse phase-HPLC as shown in chapter 3.

4.3.4.3 HMGS assays

The dependence of enzyme activity on the cofactor concentration was measured by monitoring the disappearance of acetoacetyl-CoA at 300 nm (εₘ₉₉ = 3.6) in 100 mM Tris-HCl at 30°C containing 200 μM acetyl-CoA, 15 μM acetoacetyl-CoA and 0, 1, 2, 5, or 10 mM MgCl₂. The pH dependency of AaHMGS activity was measured by the same assay by increasing the pH in 0.5-unit increments from 7 to 9 using the acetoacetyl-CoA extinction coefficients (300 nm) appropriate for each pH value. To determine the $K_m$ of acetyl-CoA, 2 μg of AaHMGS was incubated in 100 mM Tris-HCl (pH 8) at 30°C with 15 μM acetoacetyl-CoA and acetyl-CoA concentrations ranging from 4 to 400 μM, and activity was determined by monitoring the disappearance of acetoacetyl-CoA at 300 nm. The $K_m$ of acetoacetyl-CoA was determined using fixed concentration of acetyl-CoA (250 μM) and acetoacetyl-CoA concentrations ranging from 0.25 to 100 μM.
4.4 Results and discussion

4.4.1 Structural analysis of AaPMK through molecular modelling

AaPMK is a 187 amino acid protein comprised of three domains; the core region (residues 1-41, 100-107, 120-130 and 164-187), the LID region (132-164) and the substrate binding region (42-99, 108-118) (Fig. 32). It has in the center five-stranded parallel β-sheet, with a strand order of 23145, and eight α-helices (Fig. 32). The structure of AaPMK discloses the typical NMP kinase fold (Fig. 32), similar to the structure of the enzyme from Homo sapiens (HsPMK) (Chang et al., 2008). During the PMK catalysis, the P-loop or Walker A motif is involved in binding with the triphosphate, the substrate binding region binds with PM, and the LID region has an opening and closing motion, to permit the binding and release of substrates (Chang et al., 2008; Olson et al., 2009).

4.4.2. Characterization of the activity of AaPMK

A PMK has been kinetically characterized for the first time in insects. The proper expression of the functional AaPMK protein was verified by SDS-PAGE and western blot (Fig. 31). AaPMK activity was investigated at different pHs. The optimum pH for AaPMK was 7.5, which decreases in activity at pHs lower than 6.5 and higher than 8 (Fig. 33A). AaPMK shows a cation dependence on Mg$^{2+}$, with a 2 mM concentration revealing the maximal activity (Fig. 33B). Kinetic constants were determined by nonlinear regression analysis using the GraphPad Prism Software. The $K_m$ values for ATP and MA were 104 ± 11 and 190 ± 16 respectively. The $K_m$ values for PM and ADP were respectively 180 ± 39 and 107 ± 29. The $V_{max}$ was 7.9 ± 0.8 μmol/min/mg enzyme for the forward reaction, and 3.1 ± 1.2 μmol/min/mg enzyme for the backward reaction. The kinetics constants for the PMKs from the bacteria Enterococcus faecalis (EfPMK) (Doun
et al., 2005) and Streptococcus pneumonia (SpPMK) (Pilloff et al., 2003), the yeast Saccharomyces cerevisiae (SCPMK) (Garcia and Keasling, 2014) and the mammals Sus domesticus (SdPMK) (Eyzaguirre et al., 2006) and Homo sapiens (HsPMK) (Herdendorf and Miziorko, 2006) are compared with the AaPMK kinetic constants in table 5.

Addition of GPP and FPP did not affect AaPMK activity at concentrations up to 200 μM, hence feedback inhibition mechanisms can be ruled out for AaPMK.

4.4.3 Molecular model of AaFPPS

Based upon the available crystal structure of an avian, Gallus gallus FPPS (GgFPPS) (Tarshis et al., 1994), the 3-D structure of AaFPPS was modeled (Fig. 34). The homology model of AaFPPS shows that protein folded as a single domain, composed of all antiparallel α-helixes and no β-sheets (Fig. 34). Other enzymes, which also utilize isoprenyl diphosphate as their substrate, such as squalene cyclase, 5-epi-articolochene synthase, pentalenene synthase and protein farnesyl transferase, showed similar core structures formed only by α-helices (Wendt et al., 1997; Starks et al., 1997; Lesburg et al., 1997; Long et al., 1998). Hence these structures of proteins have been given the name ‘terpenoid synthase fold’ (Wang and Ohnuma, 2000).

Amino acid sequence alignments of FPPS from four insect species including Aedes aegypti (AaFPPS), Culex quinquefasciatus (CqFPPS), Anopheles gambiae (AmFPPS), and Drosophila melanogaster (DmFPPS) revealed the sequences of the five well conserved motifs typical of E-prenyltransferases (Fig. 35). The avian, Gallus gallus FPPS (GgFPPS) was also used for comparison, since its crystal structure is known (Tarshis et al., 1994) and has close similarity with the sequence of AaFPPS (44 %). The AaFPPS contains five conserved motifs found in other insects, as well as in the avian
FPPS (Fig. 8). The sequence of AaFPPS revealed two prominent aspartate rich motifs, DDXXD, known as FARM (first aspartate rich motif) and SARM (second aspartate rich motif), located in the II and V motif respectively (Fig. 35). The presence of FARM and SARM motifs represents a typical feature of E- prenyltransferases, and they have also been reported in the sequences of other enzymes that use isoprenyl diphosphate as substrate, such as monoterpen cyclases, sesquiterpene cyclases, and diterpene cyclase. The FARM motif and the first two aspartate residues in the SARM are involved in the enzyme catalytic mechanism via a metal cofactor; while the last aspartate residue is involved only in binding but not in the catalytic efficiency of the enzyme (Marrero et al., 1992; Joly et al., 1993; Song et al., 1994; Koyama et al., 1996). In addition motifs I and IV are also involved in binding with IPP, as experiments have demonstrated a very high fold increase in $K_m$ IPP of enzyme when the conserved lysine is mutated into an aliphatic amino acid in Bacillus stearothermophilus FPP synthase (Koyama et al., 1996).

**4.4.4. Partial characterization of AaFPPS activity**

The expression of a functional AaFPPS protein was verified by SDS-PAGE and western blot (Fig. 36). Optimum pH values for the FPPS previously characterized from other insects such as cotton boll weevil, Anthonomus grandis (Taban et al., 2009) and horseradish leaf beetles, Phaedon cockleariae (Frick et al., 2011) was 7.5 in 50 mM Tris-HCl buffer; hence the activity of AaFPPS was also measured under similar conditions of buffer and pH. AaFPPS yielded 84% C$_{10}$-geranyl pyrophosphate (GPP) and 16% C$_{15}$-farnesyl pyrophosphate (FPP) in the presence of Co$^{2+}$ as a cofactor; whereas it yielded 33% C$_{10}$ GPP and 67% C$_{15}$ FPP in the presence of Mg$^{2+}$ (Fig. 36). This unusual product regulation mechanism by metal cofactor was also observed in the beetle Phaedon.
cochleariae FPPS (Frick et al., 2013). The physiological significance of the product regulation of FPPS by cofactors is unknown. In A. aegypti 24 h before adult eclosion, the pool of GPP is very high (50,000 fmol/CA) and the FPP pool is low (60 fmol/CA) (Rivera-Perez et al., 2014). After the adult eclosion the size of the GPP pool decreases, and GPP becomes undetectable; while the FPP pool fluctuated from a low of 10 to a high of 100 fmol/CA, depending upon the physiological state of the CA (Rivera-Perez et al., 2014).

4.4.5 Characterization of the AaHMGS activity

The expression of the functional AaHMGS protein was verified by SDS-PAGE and western blot (Fig. 31). The activity of AaHMGS was measured using both the DNTB-CoA assay and the disappearance of acetoacetyl-CoA at 300 nm. The AaHMGS activity was investigated at different pHs. The optimum pH for AaHMGS was 8-8.5 (Fig. 37). This value is comparable to the pH 9.4, reported for the avian liver cytoplasmic HMGS, (Clinkenbeard et al., 1975), the plant Brassica juncea HMGS (pH 8.5) (Nagegowda et al., 2004), the bacterial E. faecalis HMGS (pH 9.8) (Sutherlin et al., 2002) and the archaeal Haloferax volcanii HMGS (pH, 8.5) (VanNice et al., 2013). The HMGS does not require any co-factor; however the activities of the animal recombinant enzymes were enhanced by adding Mg$^{2+}$. The AaHMGS was 180% activated in the presence of 5 mM Mg$^{2+}$, similar to that avian liver cytoplasmic HMGS (Clinkenbeard et al., 1975) and the insect, Blattella germanica (Cabona et al., 1996). On the contrary, the activity of the plant, Brassica juncea HMGS did not show any increase in the presence of metal cofactor (Nagegowda et al., 2004).
AaHMGS exhibited a $V_{\text{max}}$ of 3.3 μmol/min/mg enzyme, and the $K_m$ for acetyl-CoA and acetoacetyl-CoA were 98 μM and 14 μM respectively. These values were in the range of $V_{\text{max}}$ and the $K_m$ values reported for HMGSs of different organisms. Hymeglusin is a fungal metabolite that exhibits high specificity for inhibition of the HMGS activity (Tomoda et al., 1988). It inhibits the activity of the recombinant enzyme as well as HMGS activity in crude extracts from thorax and abdominal carcass of *A. aegypti* (Fig. 38).

4.4.6 HMGS expression and activity in the abdominal carcass of 0% Vs 20% sugar fed mosquitoes

Expression of *Aa*HMGS is found in almost all female mosquito tissues (Nouzova et al., 2011). It has been shown in various organisms that expression of HMGS transcripts can be varied with source of nutrients. In the mycelia of fungi *Ganoderma lucidum*, HMGS expression profile analysis revealed that signaling molecules such as salicylic acid, abscisic acid and methyl jasmonate up regulated *G/0*HMGS transcript levels (Ren et al., 2013). In addition carbon source has significant effects on *G/0*HMGS transcript levels (Ren et al., 2013). Similarly, expression of *Aa*HMGS is significantly reduced with silencing of TOR (Meritxell-Perez et al., 2013). In contrast we found that *Aa*HMGS transcripts do not change with the nutritional condition of the insects (Fig. 39), but its activity gets significantly reduced when mosquitoes are starved for three days (Fig. 40).

4.5 Conclusions

We have started the characterization of *Aedes aegypti* HMGS, PMK and FPPS, three additional enzymes in the JH pathway. These studies will be the foundation for
further analyses. They need to be completed in the future, but they are already revealing some interesting features that might be relevant for a better understanding of JH synthesis in mosquitoes.
Figure 28: Reaction catalyzed by HMG-CoA synthase. HMGS catalyzes the formation of HMG-CoA in three steps: acetylation, condensation, and hydrolysis (Mizioroko and Lane, 1977; Pojer et al., 2006). Step 1 is the acetylation of acetyl-CoAS to acetyl-S-enzyme intermediate, step 2 involves the condensation of acetoacetyl-CoAS and acetyl-S-enzyme to form CoAS-HMG-S-enzyme, and step 3 is the hydrolysis of CoAS-HMG-S-enzyme to produce HMG-SCoA.
Figure 29: Reaction catalyzed by PMK. *Aa*PMK catalyzes the cation-dependent reversible reaction of phosphomevalonate and ATP to form diphosphate mevalonate and ADP.
Figure 30: Reaction catalyzed by FPP synthase. FPPS catalyzes the dissociative electrophilic alkylation of IPP and DMAPP or GPP, which is divided into three steps. In step 1, the C1–oxygen bond in DMAPP or GPP ruptures to generate a resonance stabilized allylic cation. In step 2, this allylic cation alkylates the double bond in IPP to produce a tertiary carbocation. During step 3, hydrogen is subsequently eliminated from C2 of the IPP unit to produce a new allylic diphosphate which is one isoprene unit longer than the substrate (Poulter and Rilling, 1978; Poulter, 2006).

\[ \text{R} = \text{CH}_3, \text{C}_6\text{H}_{11} \]
Figure 31: SDS-PAGE analysis of the recombinant mosquito *Aa*HMGS, *Aa*PMK and *Aa*FPPS. Lanes are: 1, molecular standard; 2, *Aa*HMGS; 3, *Aa*PMK; 4, *Aa*FPPS. The molecular weights of the protein standard are depicted on the Y axis.
Figure 32: Homology model of the overall fold of *Aa*PMK. The core region is shown in cyan, substrate binding region is shown in green, LID region is shown in yellow, Walker A loop motif or P-loop is shown in red, and Walker B motif is shown in pink.
Figure 33: Characterization of AaPMK activity. The kinase activity was measured by an enzyme coupled spectrophotometric assay; ATP and PM were held constant, and data were normalized to the maximum observed reaction velocities. To ensure that PMK was the rate-limiting enzyme, when necessary the results were verified with the following conditions: doubling the amount of PMK added doubled the observed rate, doubling the amount of PK and LDH did not affect the observed rate, and doubling the PEP concentration did not affect the observed rate. (A) pH curve. Two different buffers were used: MES at pH 6 to 7 and Tris-HCl at pH 7 to 9. (B) Effect of metal cofactors Mg$^{2+}$ was assayed for the AaPMK activity. Each value represents the means ± S.E. of three replicate assays. Relative activity is defined as a percentage of the highest value recorded.
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<th>Family</th>
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<th>$AT$</th>
<th>$K_m$</th>
<th>$PH$</th>
<th>$K_m$</th>
<th>$ADI$</th>
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<td>NMP</td>
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<td>104 ± 11</td>
<td>190 ± 16</td>
<td>107 ± 29</td>
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<td>52 ± 1</td>
<td>34 ± 3</td>
<td>47 ± 5</td>
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<td>350 ± 20</td>
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Table 5: Kinetics of PMK. *AaPMK* kinetics were compared with the kinetics of *HsPMK* (*Homo sapiens* PMK), *SdPMK* (*Sus domesticus* PMK), *ScPMK* (*Saccharomyces cerevisiae* PMK), *SpPMK* (*Streptococcus pneumoniae* PMK), *EfPMK* (*Enterococcus faecalis* PMK).
Figure 34: Homology model for the overall fold of *AaFPPS*. Three dimensional structure of *AaFPPS*. Motifs are indicated by colors; motif I (red), motif II (blue), motif III (yellow), motif IV (purple), motif V (Orange). The structure was constructed by PyMOL using the avian FPPS (PDB: 1ubx.1) as template.
Figure 35: Sequence alignment of the deduced amino acid sequence from insect FPPSs. The sequence of AaFPPS was aligned with insect FPPS from Culex quinquefasciatus (CqFPPS), Anopheles gambiae (AgFPPS), Drosophila melanogaster (DmFPPS), and the avian Gallus gallus (GgFPPS). Identical residues are indicated by an asterisk. The box indicates five highly conserved regions. Accession number AaFPPS (AAEL003497), AgFPPS (AGAP007104), CqFPPS (EDS25700.1), DmFPPS (NP_477380.1), GgFPPS (1UBV_A)
Figure 36: Effect of metal cofactors on enzyme activity and product formation of AaFPPS. 100 μM IPP and 100 μM DMAPP were incubated in presence of 5 mM Mg^{2+} (A) or 1 Mm Co^{2+} (B). Reactions were started with the addition of 4 μg recombinant enzyme. Formation of FPP and GPP were measured as described in method section.
Figure 37: Characterization of *Aa*HMGS activity A) Magnesium dose-dependent increases of activities. B) Optimum pH determinations. Each value represents the means ± S.E.M. of three replicate assays. Relative activity is defined as a percentage of the highest value recorded.
Figure 38: Inhibition of *Aa*HMGS abdominal carcass activity by hymeglusin. The reaction was performed by the DNTB-CoA assay. 1 mg of protein was added in 24 h 20% SF females after adult eclosion in the presence and absence of hymeglusin. Each point (± SEM) represents the mean of 3 biological replicates of extracts from 5 AC from female mosquitoes. Significant differences (p<0.01) were determined with one way ANOVA followed by Tukey’s test.
Figure 39: Activity of HMGS from female AC extracts. The reaction was performed by the DNTB-CoA assay. 1 mg of protein was added in each of the three points; 24 h 20% SF, 96 h 20% SF and 96 h 0% SF females after adult eclosion. Each point (± SEM) represents the mean of 3 biological replicates of extracts from 5 AC from female mosquitoes. Significant differences (p<0.05) were determined with one way ANOVA followed by Tukey’s test.
Figure 40: Expression of HMGS mRNAs from female abdominal carcass (AC) extracts. Expression of HMGS AC mRNAs of 24 h 20% SF, 96 h 20% SF and 96 h 0% SF females after adult eclosion. HMGS mRNAs are expressed as copy number of HMGS mRNA/10,000 copies of rpl32 mRNA. Each RT-PCR data point is average of three independent biological replicates of 5 AC. Significant differences (p<0.05) were determined with one way ANOVA followed by Tukey’s test.
4.6 References


Chapter 5: Conclusions and future directions

5.1 Conclusions

This dissertation is a comprehensive study of five *Aedes aegypti* CA enzymes, HMGS, MK, PMK, FPPS, and FPPase involved in JH synthesis. The study mainly focused on the expression and characterization of recombinant protein, the analysis of their kinetics and inhibition constants, as well as the understanding the importance of these enzymes in the control of JH biosynthesis rates.

In chapter 2 FPP phosphatase responsible for the transformation of FPP into FOL in the CA of *A. aegypti* was characterized at molecular and biochemical levels. Some of the observations from chapter 2 include:

1. Identification of *AaFPPase*-1 and *AaFPPase*-2 as members of the NagD family of the Class IIA C2 cap-containing haloalkanoic acid dehalogenase (HAD) super family that efficiently hydrolyzed FPP and GPP but not IPP.
2. Different FPPase activities in CA of sugar-fed as compared to blood-fed females.
3. Injection of dsRNAs resulted in a significant reduction of both *AaFPPase*-1 and *AaFPPase*-2 mRNAs, but only former resulted in a significant decrease of JH biosynthesis.
4. *AaFPPase*-1 appears to be the major enzyme involved in the catalysis of FPP into FOL in the CA of *A. aegypti*.
5. N-acetyl-S-geranylgeranyl-L-cysteine appears as a potent inhibitor of *AaFPPase* 1 and *AaFPPase* 2 in blocking JH synthesis in *A. aegypti*.

Chapter 3 describes the first characterization of a mevalonate kinase in insects.
Some of the conclusions of the studies included are:

1. *AaMK* is a class I MK that has the typical structure and functional features of other members of the GHMP kinase family.

2. Recombinant *AaMK* is found to be inhibited by isoprenoids; with a pattern similar to those of other eukaryotic MKs.

3. Inhibition of the activity of MKs by isoprenoids is found to be competitive with respect to ATP and the eukaryotic enzymes are 1000-fold more sensitive to the inhibition by isoprenoids than bacterial and archaeal enzymes.

4. The MK activity in thoraces extracts of sugar-fed (SF) mosquitoes was also significantly reduced in the presence of FPP.

Chapter 4 reports the partial characterization of the activities of recombinant HMGS, PMK and FPPS from *A. aegypti*. Some of the conclusions of the studies included are:

1. HMGS transcripts do not change with the nutritional condition of the insects. However its activity was reduced when mosquitoes are starved for three days.

2. Hymeglusin is a specific inhibitor of HMGS that could be used to block the mevalonate pathway in *A. aegypti*.

3. *AaPMK* is a member of the nucleoside monophosphate family.

4. The present study is the first report on kinetics of *AaPMK* for both forward and backward reactions in insects. *AaPMK* catalytic activity is not inhibited by any of the downstream metabolites.

5. Recombinant *AaFPPS* possesses an interesting product regulation mechanism; it alters the chain length of its products depending on the cofactor present.
5.2. Future directions

All the thirteen corpora allata enzymes involved in the pathway of juvenile hormone biosynthesis have been identified in the mosquito *A. aegypti*. With the inclusion of appropriate cofactors, CA extracts are capable of *de novo* synthesis of JH metabolites from precursors (Rivera-Perez *et al*., 2013, 2014; Nyati *et al*., 2013); these in vitro assay have been used to study the endogenous activities of eight of these enzymes, HMGS, MK, PMK, FPPS, FPPase, FOLD, FALD and JHAMT. These eight enzymes along with IPPI have been expressed as recombinant proteins and their activities were characterized in vitro (Mayoral *et al*., 2009a, 2009b; Diaz *et al*., 2012; Rivera-Perez *et al*., 2013; Nyati *et al*., 2013). Only four enzymes remain to be characterized in mosquitoes, acetoacetyl-CoA thiolase, HMGR, MDD and epoxidase.

The enzymes of the JH-branch have been considered as more suitable for insecticides target of JH biosynthesis; given that inhibitors targeting them are less likely to affect non-insect organisms. Inhibitors directed at the MVP enzymes may lack insect-specificity; however, in some groups of insects, MVP enzymes may display unique features that will permit the development of target-specific inhibitors (Cousson *et al*., 2013). There are examples of such effective inhibitors; 6-fluoromevalonate 5-diphosphate fluoromevalonolactone (FMev) completely inhibits the activity of lepidopteran MDD (Quistad *et al*., 1981; Baker *et al*., 1986; Cusson *et al*., 2013). Inhibitors of FPPSs, such as lipophilic bisphosphonates, have been designed as drugs and herbicides (Oldfield, 2010; Cromartie, 1991). Lipophilic bisphosphonates effectively inhibit protein prenylation and invasiveness in tumor cells and has been used as anticancer agents (Zhang *et al*., 2009; Oldfield, 2010). The completion of the functional analyses of all the
enzymes involved in the biosynthesis of JH in *A. aegypti* will permit the search for specific inhibitors that target some of these enzymes and disrupt the mosquito reproductive physiology.

Recombinant enzymes have been used in building of wide range of natural and synthetic isoprenoids to be used as medicines, cosmetics, flavors, fragrances and biofuels (Hale *et al*., 2007; Lacaze *et al*., 2011; Peralta-Yahya *et al*., 2011; Zhang *et al*., 2011). Recent attempts include the production of isoprenoid-based biofuels that might help to overcome rising petroleum costs, trade imbalances and environmental concerns (Peralta-Yahya *et al*., 2011; Zhang *et al*., 2011). Artemisinin, an antimalarial drug has been enzymatically synthesized from acetyl-CoA using genetic engineering at very low costs (Hale *et al*., 2007; Lacaze *et al*., 2011). Similar approaches could be applied for the development of a multienzyme JH pathway bioreactor. The enzymes of the JH pathway may be segregated in a module, immobilized on a membrane, or attached to a support for the development of the JH bioreactor. This could result in significant enhancements in yield, purity, production time and cost when compared to traditional chemical synthetic methods for the synthesis of JHs, pheromones, defensive secretions and many other isoprenoid-derived compounds. Additionally, linking the bioreactor with the HPLC-fluorescent detection assay could help in the study of metabolites flux and changes in pool sizes, helping to model JH regulation, as well as the design of a multi-enzyme inhibitor.
5.3 References


7. Nyati P, Rivera-Perez C, Noriega FG Negative feedback by isoprenoids upon mevalonate kinase activity might regulate juvenile hormone synthesis in *Aedes aegypti*. In prep


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