University of Alberta

Mapping and characterization of a caffeine-sensitive mutant in Drosophila melanogaster

by

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in

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Abstract

The caffeine-sensitive *Drosophila melanogaster* mutant $huc95^{DE}$ was previously isolated in a screen for hydroxyurea and/or caffeine sensitivity and mapped to 95D11-E7, delineating $huc95^{DE}$ to 23 possible genes. I have mapped $huc95^{DE}$ to 95E1-E7-E8, narrowing the number of candidates to 9, resulting in a renaming to $huc95^{E}$. When reared on caffeine, $huc95^{E}$ exhibit small, disorganized eyes with fewer ommatidia that appear irregular and fused when homozygously expressed in the eye. $huc95^{E}$ hemizygotes exhibit pupal lethality at caffeine concentrations as low as 0.25 mM, but can survive in normal media suggesting that $huc95^{E}$ is not required for viability. In the presence of caffeine, $huc95^{E}$ hemizygotic third instar larvae exhibit degenerating imaginal discs and hemizygotic pupae contain no visible adult structures. Staining of these discs with acridine-orange and caspase-3 demonstrates high levels of apoptosis, providing and explanation for these developmental defects.

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List of Symbols and Abbreviations

	Male		Virgin female
AAA	ATPase Associated with Various Cellular Activities	AC	Adenylyl Cyclase
AIF	Apoptosis-Inducing Factor	AMP	Adenosine Monophosphate
Apaf-1	Apoptosis Protease Activation Factor-1	ATM	Ataxia Telangiectasia- Mutated
ATP	Adenosine triphosphate	ATR	ATM and Rad3-related
BrdU	Bromodeoxyuridine	ca	Claret
Ca ²⁺	Calcium Ions	cAMP	Cyclic AMP
CDK1	Cyclin Dependent Kinase-1	C Nap-1	Centrosomal Nek2- Associated Protein-1
СуО	Second chromosomal balancer	DAG	Diacylglycerol
Df	Deficiency	е	Ebony
EGUF	Eyeless promoter-GAL4; UAS Flp-recombinase	Elav	Embryonic Lethal Abnormal Visual
EMS	Ethyl Methanosulphonate	ER	Endoplasmic Reticulum
FMW	First Mitotic Wave	GFP	Green Fluorescent Protein
GMR-hid	<i>Glass Multiple Reporter</i> promoter- <i>Head</i> <i>Involution Defective</i>	GPCR	G-Protein Coupled Receptor
G Protein	GTP/GDP binding Protein	GTP/GDP	Guanosine Triphosphate /Diphosphate
Gy	Gray	HU	Hydroxyurea
Hu	Humoral	huc	Hydroxyurea and/or Caffeine Sensitive
IAP	Inhibitor of Apoptosis Protein	IQ	Isoleucine-Glutamine motifs
IP ₃	Inositol 1,4,5 Trisphosphate	IP ₃ R	IP ₃ Receptor

LC3	Microtubule Associated Protein-1 Light Chain 3	mМ	Millimolar
Nek	NIMA Related Kinase	NIMA	Never In Mitosis in Aspergillus nidulans
p70S6K	p70 Ribosomal S6 Kinase	PAS	Preautophagosome
PH-3	Phosphohistone-3	PI3K	Phosphoinositide 3 kinase
РКА	Protein Kinase A	РКВ	Protein Kinase B
РКС	Protein Kinase C	RyR	Ryanodine Receptor
Sb	Stubble	Ser	Serrate
SMAC	Second Mitochondrial Activator of Cell Death	SMW	second mitotic wave
Tb	Tubby	<i>TM6</i>	Third chromosome balancer
ТМЗ	Third chromosome balancer	TP	Transposition
UAS	Upstream Activating Sequence	Ubx	Ultrabithorax
w	White		

Chapter 1: Introduction

Caffeine is the most widely used psychoactive drug in the world today, with total consumption levels on average of 76 mg/person/day, reaching 210-238 mg/person/day in the United States and Canada, and up to 400 mg/person/day in the United Kingdom and Scandinavian countries (Gilbert, 1984; Barone and Roberts, 1996; Mandel, 2002). Much of this usage (which transcends age and cultural boundaries) occurs because caffeine acts as a neurological stimulant to suppress mental and physical fatigue and improve attention and arousal, without substantial or clearly documented negative side effects at typical consumption levels. Caffeine also exhibits a wide variety of other pharmacological effects, including sensitizing cells to irradiation and genotoxic drugs (presumably by overriding cell cycle checkpoint responses), promoting apoptosis, and altering cellular calcium levels through its effects on calcium channels (Lau and Pardee, 1982; Schlegel and Pardee, 1986; He et al., 2003; Zucchi and Ronca-Testoni, 1997; Rousseau et al., 1988). These additional effects have made caffeine a useful tool in research for studying cellular proliferation, cell death, and molecular mechanisms involving calcium signalling including muscle contraction or calcium channel excitation. Despite its wide consumption and use as a reagent in research, the mechanism underlying these diverse actions of caffeine have not been fully elucidated. Therefore, it is important (both from a practical research standpoint as well as medically and clinically) to understand the mechanisms underlying the pharmacological activities of caffeine. My project represents an effort to characterize gene functions that protect cells from caffeine toxicity in Drosophila.

Section 1-2: Caffeine and adenosine

Many of caffeine's effects are thought to stem from structural similarities to the nucleoside adenosine and other structurally related derivatives (such as 2-deoxyadenosine, adenine, and the signalling molecule, cyclic adenosine monophosphate (cAMP) - see Figure 1-1) that are involved in a multitude of signalling pathways regulating various cellular and physiological processes (Nishijima *et al.*, 2003).



Figure 1-1. Comparison of the structures of caffeine, adenosine and cyclic AMP

Concerning adenosine, it is a common metabolite made by all cells and is often formed as a by-product of energy utilization through the catabolism of adenosine triphosphate (ATP). Levels of adenosine play a central role in the energy metabolism of a cell by acting as a key indicator of energy utilization, initiating certain processes when imbalances between energy consumption and energy supply occur (Dunwiddie and Fredholm, 1988). Consequently, altering levels of adenosine can modulate many diverse cellular processes including cell proliferation and cell death, and can physiologically modulate changes in the cardiovascular system, central nervous system, gastrointestinal tract and immune system, as well as mast cell degranulation, pain modulation, and asthma (Schulte and Fredholm, 2003). Because of these effects, adenosine levels must be constantly regulated to prevent untimely events or processes from occurring (Jacobson *et al.*, 1999; Merighi *et al.*, 2003). As a result, constant levels of adenosine are maintained by the phosphorylation or deamination to AMP or inosine respectively, by the enzymes adenosine deaminase or adenosine kinase.

Because of its similarity to adenosine, caffeine is thought to exert its effects by interacting with adenosine receptors or other interacting partner proteins and either mimicking or antagonizing adenosine. In fact, it was the study of the antagonistic effects of caffeine that first indicated the existence of adenosine receptors (De Gubareff and Sleator Jr., 1965). Adenosine receptors are subdivided into four different receptor subtypes (classified as A1, A2A, A2B, and A3, respectively), originally classified according to their abilities to either inhibit (as is the case of A_1 receptors) or stimulate (as is the case of A₂ receptors) levels of the ubiquitous secondary messenger, cyclic AMP (cAMP) (van Calker et al., 1978; van Calker et al., 1979; Londos et al., 1980; Fredholm, 1982). Stimulatory A₂ receptors were further subdivided according to their abilities to stimulate cAMP production within brain slices at low (A_{2A}) and high (A_{2B}) adenosine concentrations, which was further supported through ligand-binding assays and molecular biological analysis revealing differential affinities to adenosine derivatives (Daly et al., 1983; Bruns et al., 1986; Fredholm et al., 2001). A₃ receptors (which generally result in a decrease in cAMP levels) were discovered on the basis of sequence similarity during the molecular cloning of a rat testis cDNA library and were classified because of their insensitivity to the antagonistic actions of methylxanthines, a trait which was responsible for this class remaining unknown until 1992 (Zhou et al., 1992; Palmer et al., 1995). All

adenosine receptors belong to a class of transmembrane receptors that are associated with guanosine triphosphate/diphosphate (GTP/GDP)-binding proteins (G-proteins) called Gprotein coupled receptors (GPCRs). G-proteins consist of a heterotrimeric complex of three subunits (labelled α , β , and γ , respectively) that contain a binding pocket for GTP or GDP within the first (α) subunit (Gilman, 1987) as shown in Figure 1-2A. Each class of adenosine receptor associated with particular types of different G-proteins that are thought to impart selective activities for each receptor, possibly serving as a link to various other cell signalling pathways (Schulte and Fredholm, 2003). For instance, A1 and A_3 receptors have been found to interact with $G_{11/2/3}$, and G_0 proteins (causing decreased levels of cAMP), whereas A_{2A} and A_{2B} receptors have been found to associate with G_S, G_{OLF}, and G_{15/16} receptors, and G_S, G_{O/11} proteins, respectively (resulting in increased levels of cAMP) (Schulte and Fredholm, 2003). A₃ receptors have also been found associated with $G_{Q/11}$ as well, suggesting that these receptors could serve dual functions of either stimulating or inhibiting cAMP production, depending upon which complex is formed. As a result, the interaction of adenosine receptors with various different G-proteins further adds to the complexity of the regulation and signalling of these receptors and consequently, the possible effects that adenosine analogs such as caffeine may exert.

Section 1-3: Adenosine and cAMP signalling

Much of adenosine's ability to elicit effects on a multitude of systems involves the formation of the ubiquitous secondary messenger cAMP. cAMP is produced in response to the availability of nutrients, or the presence of hormones, ions or other



Figure 1-2. Mechanism of adenosine receptor stimulation and the consequent activation of cAMP production. In its unstimulated state (A), the transmembrane G-protein coupled receptor (GPCR) contains GDP in a binding pocket of the first α subunit in a heterotrimeric complex associated with the receptor. When stimulated by a ligand (B), GDP is replaced by GTP, which signals dissociation of the $\beta\gamma$ subunits and consequent activation of adenylyl cyclase by activated G α , which converts ATP to cAMP.

signalling molecules (Daniel et al., 1998). cAMP transduces signals from activated GPCRs by activating protein kinase A (PKA), which then initiates the response by phosphorylating its substrates (Daniel et al., 1998). Upon stimulation of GPCRs by a specific ligand, an allosteric change within the α subunit of the G-protein in the receptor causes the G-protein to become active by the replacement of GDP for activating GTP (Figure 1-2B). This causes the α subunit to become activated, dissociating from the β and γ subunits (still held together as a dimer) and activating the enzyme adenylyl cyclase (AC) (Daniel et al., 1998). AC catalyzes the formation of cAMP from ATP, which then interacts with the effector molecule PKA: the apical kinase in the cAMP-signalling cascade. PKA exists as a tetrameric holoenzyme composed of two catalytic subunits and a homodimer of regulatory subunits. Upon activation by cAMP at the regulatory domain, the catalytic subunits dissociate from the holoenzyme and phosphorylate serine (S) or threonine (T) residues in a specific RRX(S/T) motif (whereby R represents arginine and X represents any amino acid residue), regulating a variety of intracellular proteins, enzymes, ion channels, and gene expression through direct interaction with such transcription factors as cAMP response element binding proteins (CREB), cAMP response element modulator (CREM), activating transcription factor-1 (ATF-1), NF-KB, and nuclear receptors (Figure 1-3) (Daniel et al., 1998). As a result, many different systems and processes can be regulated through cAMP signalling, which in turn, can be activated by stimulation of adenosine receptors (Della Fazia et al., 1997). Levels of cAMP signalling are controlled by the activity of phosphodiesterases, (which serve to break down intracellular cAMP, reducing signal by reducing the secondary messenger), phosphatases (which reduce signalling by counteracting the kinase activity of PKA), and



Figure 1-3. The cAMP pathway. Upon formation of cAMP from ATP by adenylyl cyclase, this secondary messenger molecule can interact with the effector molecule PKA at regulatory (R) domains to free the catalytic subunits (C) which translocate into the nucleus and serve to phosphorylate transcription factors and enzymes such as cAMP response element binding proteins (CREB) and cAMP response element modulators (CREM). These factors can then interact with cAMP response elements (CRE) to bring about regulation in various cellular processes. (Adapted from Della Fazia *et al.*, 1997)

PKA inhibitors (which prevent the phosphorylation of important substrates through inhibition of PKA itself) (Daniel *et al.*, 1998). Caffeine can interfere with cAMP signalling and affect cellular processes regulated by this pathway by its agonistic or antagonistic actions towards different adenosine receptors, or by its actions (at concentrations in the mM range) as a phosphodiesterase inhibitor, preventing breakdown of cAMP (Figure 1-4) (Vernikos-Danellis and Harris III, 1968). As a result, the mechanisms of action of caffeine are not yet clearly defined, since various steps can be affected.



Figure 1-4. The generation and removal of cAMP. The signalling molecule cAMP is synthesized by the enzyme adenylyl cyclase from ATP upon activation from GPCRs. To regulate intracellular levels, cAMP is hydrolyzed by cAMP phosphodiesterase to form AMP. This process is inhibited by caffeine and may result in some of the effects mediated by this drug.

Section 1-4: Caffeine and calcium signalling

Caffeine affects calcium levels by influencing the regulation of ion channels and adenylyl cyclase signalling pathways through inhibition of adenosine receptors. Normally, the amount of free intracellular calcium ions (Ca^{2+}) is tightly regulated such that the concentration of extracellular Ca^{2+} is about four orders of magnitude higher (approximately 1.2 mM) than that within the cell (approximately 100 nM in resting cells) (Hanson et al., 2004; Saris and Carafoli, 2005; Waring, 2005). This is accomplished by controlling the entry or efflux of Ca^{2+} , and the active sequestration and storage of intracellular Ca²⁺ within intracellular compartments (such as the endoplasmic reticulum (ER) or mitochondria) through various channels and transporters. Increased levels of cvtosolic Ca²⁺ are made available through the controlled release from ER pools, where they can interact with molecules that require Ca^{2+} binding for regulation in various biochemical and signal transduction pathways, or act by themselves to signal other events in the cell. This regulation allows Ca^{2+} to act as a general signalling molecule for many different cellular, physiological and developmental events, since the release or entry of only a small amount of Ca^{2+} can significantly alter the concentration gradient between intra and extracellular Ca²⁺. Many extracellular signals such as hormones, neurotransmitters, and growth factors, as well as membrane depolarization or mechanical and environmental stress, initiate their respective cellular responses by a regulated release of Ca^{2+} within the cell. With regard to adenosine receptors, certain G-proteins of the G_0 subtype can activate phospholipase C, an enzyme involved in the hydrolysis of phospholipids on the inner layer of the plasma membrane (Yakel et al., 1993; Gao et al., 1999; Linden et al., 1999). When phospholipase C is activated, it hydrolyzes

phosphatidylinositol 4,5-bisphosphate (PIP₂), producing the secondary messengers diacylglycerol (DAG) and inositol 1,4,5-trisphosphate (IP₃) (see Figure 1-5). DAG remains in the inner layer of the plasma membrane and recruits protein kinase C (PKC), a calcium dependent protein kinase responsible for phosphorylating various proteins to bring about the associated changes in the cell, while IP₃ diffuses and binds to IP₃



Figure 1-5. Calcium signalling mediated by phospholipase C. The intracellular concentration of calcium (Ca²⁺) is tightly controlled such that its release from Ca²⁺ stores such as the endoplasmic reticulum (ER) can be used to signal various processes through activation of a Ca²⁺ dependent general protein kinase PKC. Phospholipase C (which may be activated through adenosine signalling) hydrolyzes phosphatidylinositol 4,5-bisphosphate (PIP₂) to produce the secondary messengers inositol 1,4,5 trisphosphate (IP₃) and diacylglycerol (DAG). DAG recruits PKC, while IP₃ signals the release of Ca²⁺ from the ER for PKC activation.

receptors (IP₃R) in the ER (the main storage and release organelle of intracellular Ca²⁺), causing a release of Ca^{2+} within the cytosol. IP₃R are large (~1200 kDa) tetrameric protein whose subunits form a Ca²⁺ channel (through interaction of integral membranespanning carboxy-terminal domains) with an exposed cytoplasmic amino-terminal domain responsible for binding IP₃ (Hanson et al., 2004) Activation of IP₃R however, is controlled not only by binding of IP₃, but also by the binding of free Ca^{2+} (in amounts of up to 500 nM) to an exposed cytoplasmic amino acid stretch between the carboxy and amino-terminal domains (Hanson *et al.*, 2004). The binding of Ca^{2+} at this concentration acts synergistically with IP₃ to induce the opening of the channel and release of Ca²⁺. At concentrations higher than 500 nM however, IP₃R opening is inhibited, serving as a control mechanism to prevent pathological Ca^{2+} increases in the cytosol. Because of the similarity of caffeine and adenosine, interaction of caffeine at these receptors could affect downstream regulation of Ca^{2+} release in certain situations. In addition to this, another class of Ca²⁺ receptors with sequence and structural similarities to IP₃R are also present within the ER, called ryanodine receptors (RyR). RyR were named because of their sensitivity to the plant alkaloid ryanodine, and have also been implicated in the regulation of cytosolic Ca²⁺ (Mignery et al., 1989). RyR are also large (approximately 2.3 Mda) tetrameric proteins, but differ from IP₃R in their biophysical and pharmacological properties. For example, the activity of RyR is increased by treatment with negatively charged polyanionic molecules (such as pentosan polysulphate, polyvinyl sulphate and heparin), whereas the activity of IP₃R is decreased (Bezprozvanny et al., 1993; Palade et al., 1989). Furthermore, treatment with protamine (a positively charged molecule) could reversibly inhibit the activity of RyR (Koulen and Erlich, 2000). These studies suggest

that the high negative charge of these polyanions served to increase the local negative charge of RyR, attracting more Ca^{2+} to the receptor thereby activating it, and that the positive charge of protamine indirectly blocks the channel by creating a cation-repelling environment at the pore complex at sites of Ca^{2+} regulation. It has also been reported that RyRs are sensitive to treatment with caffeine, evoking Ca^{2+} release (consequently, caffeine is often used in studies involving these receptors) (Rousseau *et al.*, 1988; McPherson *et al.*, 1991). As a result, increases in the levels of Ca^{2+} in the cytosol (possibly by the action of caffeine on RyR) could serve to also affect other channels, thereby further increasing the cytosolic concentration.

Section 1-5: Excess calcium levels can signal apoptosis

One important means of regulating cytosolic calcium levels and coordinately regulating respiration is by controlling the intake of Ca²⁺ into the mitochondria (the site for aerobic respiration and generation of ATP). Elevations of calcium levels within the cytosol of metabolically active cells can signal increases in the activity of citric acid cycle enzymes within the mitochondrial matrix, resulting in an increase in the rate of ATP production (Hanson *et al.*, 2004; Saris and Carafoli, 2005). In addition to providing the necessary energy requirements to carry out basic cellular metabolism, this increased rate of ATP production can also be used to re-establish the Ca²⁺ gradient through the activation of Ca²⁺ ATPases (ATP-dependent Ca²⁺ pumps that actively sequester Ca²⁺ back into storage organelles, or expel Ca²⁺ from the cell, and are crucial for keeping the concentration of intracellular Ca²⁺ at approximately 100 nM). If the concentration of Ca²⁺ becomes too high within the mitochondrial matrix however, the mitochondria could swell, rupture, or undergo permeability changes, which may release pro-apoptotic proteins into the

cytoplasm. These pro-apoptotic proteins include cytochrome C, second mitochondrial activator of cell death (SMAC/Diablo), and apoptosis-inducing factor (AIF) (Hanson et al., 2004). Cytochrome C is a 13 kDa nuclear encoded protein that serves as an electron carrier within the electron transport chain and is important for the establishment of the electropotential gradient in the generation of ATP through oxidative phosphorylation. Release of cytochrome C into the cytoplasm acts as a general signal for the initiation of apoptotic events however, by interacting with pro-apoptotic proteins to form the apoptosis promoting structure called the apoptosome, described below (Wang et al., 2005). SMAC/Diablo commits the cell to programmed cell death by interacting with, and suppressing the actions of inhibitors of apoptosis proteins (IAP) which negatively regulate apoptosis by binding key apoptotic proteins such as caspase 9 (Wang et al., 2005; Du et al., 2000). AIF is a nuclear encoded flavoprotein that resides in the mitochondrial intermembrane space, and is responsible for the condensation and fragmentation of nuclear DNA upon its release into the cytoplasm, an event that is associated with apoptosis (Susin et al., 1999; Wang et al., 2005). The process of apoptosis is mediated by the activation of caspases (aspartate-directed cysteine proteases that cleave specific substrates at cysteine residues) from inactive zymogens (proenzymes). These activated caspases then serve as initiators and effectors of programmed cell death (see Figure 1-6). Caspases are expressed as single chain proenzymes containing an NH2-terminal domain, a large (20 kDa) catalytic subunit and a smaller (10 kDa) subunit (Wang et al., 2005). Caspases can be classified into two categories. Initiator caspases such as caspase 2, 8, 9, and 10, are characterized by long N-terminal prodomains that contain conserved protein-protein interaction motifs.

Effector caspases such as caspase 3, 6, and 7, possess either smaller, or no prodomains at all. Upon proteolytic activation, initiator caspases recruit and activate effector caspases, which cleave their target proteins to bring about the orderly demise of the cell. This process is initiated when the released cytochrome C and other components such as dATP, apoptosis protease activation factor-1 (Apaf-1), and pro-caspase 9 assemble into a high molecular weight heptameric wheel-like complex called the apoptosome, consisting of seven Apaf-1/caspase 9 dimers that transiently associate with cytochrome C (Wang et al., 2005). The formation of the apoptosome allows for the efficient activation of initiator caspase 9 as the association of cytochrome C to pro-caspase 9 allows Apaf-1 to interact with pro-caspase 9 at a caspase-associated recruitment domain (CARD) in the N terminus, resulting in its activation. Once activated, caspase 9 cleaves effector caspases 3, 6, and 7, which, upon activation, will cleave at aspartate-cysteine residues on intracellular substrates such as the DNA repair enzyme poly(ADP-ribose) polymerase (PARP) (which when cleaved, translocates AIF from the mitochondria to the nucleus), caspase-activated **D**Nase (CAD), nuclear lamins, gelsolin (an actin-capping protein important in maintaining cell shape and structure in the transition of gel to sol states of cytoplasm), and fodrin (a protein important in maintaining cell shape, stabilizing membrane structures, and linking the cytoskeleton to plasma membranes or intracellular vesicles). All of these processes serve to destabilize nuclear and cytoplasmic integrity in a characteristic sequence of apoptotic events.

The cycling of Ca^{2+} between the ER and mitochondria is a key event in the signalling of apoptosis with the process of depleting ER stores emerging as an important trigger. Ca^{2+} continuously cycles between the mitochondria and ER, allowing



Figure 1-6. Mechanism of calcium-induced apoptosis. Mitochondrial rupture mediated by excessive Ca^{2+} influx can result in the release of cytochrome C, <u>apoptosis</u>inducing <u>factor</u> (AIF), and Diablo (*aka* <u>second</u> <u>mitochondrial</u> <u>activator</u> of <u>c</u>ell death (SMAC)) into the cytoplasm to initiate apoptosis. Cytochrome C interacts with <u>apoptosis</u> <u>protease</u> <u>activation</u> <u>factor</u> 1 (Apaf-1) and pro-caspase 9 to form the apoptosome, which activates caspase 9 to induce cleavage of effector caspases 3, 6, and 7, which directly bring about the events of apoptosis. AIF translocates into the nucleus (which is mediated by the activity of the activated caspases) to bring about DNA fragmentation while Diablo commits the cell to programmed cell death by inhibiting inhibitors of <u>apoptosis</u> <u>proteins</u> (IAP) that normally prevent apoptosis from occurring.

mitochondria to regulate Ca^{2+} signalling and levels within the ER (Demaurex and Distelhorst, 2003). Depletion of Ca^{2+} from the ER pool, and the consequent increase of Ca^{2+} in mitochondria, signals apoptosis and can provide some insight into how caffeine might induce apoptosis. For instance, Chinese hamster ovary cells transfected with the RyR and treated with 10 µM of ryanodine (which is just enough to lock the RyR in an open state) and 10 mM caffeine, undergo apoptosis caused by the rapid caffeine-induced depletion of Ca^{2+} from the ER (Pan *et al.*, 2000). Addition of caffeine has been linked to apoptosis in a variety of cell types, although the dosage required for this effect differs amongst different cell types and the role that caffeine plays in these apoptotic events might be more complex than interaction at RyRs (Fernandez *et al.*, 2003; Jafari and Rabbani, 2000; He *et al.*, 2003). For instance, evidence in JB6C141 cells suggests that caffeine-induced apoptosis is mediated by the tumor suppressor p53, which activates the proapoptotic protein Bax to release cytochrome C (He *et al.*, 2003). This ultimately results in the activation of caspase 3 (the major effector caspase responsible for apoptosis) to mediate apoptotic events.

Section 1-6: Caffeine can inhibit signalling by phosphoinositide 3-kinase

Caffeine can affect the signalling molecule phosphoinositide (PI) 3-Kinase (PI3K), which has been implicated in such processes as cellular proliferation, apoptosis, differentiation, and metabolic regulation (Foukas *et al.*, 2002; Katso, *et al.*, 2001; Osaki *et al.*, 2004; Franke *et al.*, 2003). PI3Ks phosphorylate the D3'-OH position of the inositol ring of phosphoinositides to create secondary lipid messengers which interact with other signalling molecules downstream to exert their effect. They can be further subdivided into four classes (classes I_A , I_B , II, and III, respectively) based upon lipid

substrate specificity, structure, and mode of regulation (Vanhaesebroeck et al., 1997; Fruman et al., 1998). Class I enzymes are the most well studied class of PI3Ks and consist of a 110 kDa catalytic subunit (of either α , β , δ , or γ) associated with a regulatory adaptor subunit (p85 α , p85 β , p55 α , p55 γ , p50 α , p87, or p101) linking the enzyme to signalling molecules upstream (Fruman et al., 1998). Upon activation, class I enzymes phosphorylate phosphatidylinositol 4,5-bisphosphate (PtdIns-4,5-P₂) to produce the secondary lipid messenger PtdIns-3,4,5-P₃. These lipid messengers interact with other signalling molecules such as protein kinase B (PKB/Akt) and p70 ribosomal S6 kinase (p70S6K) at <u>p</u>leckstrin <u>h</u>omology (PH) domains to mediate their effects. This class is further subdivided into classes I_A and I_B, with I_A consisting of PI3Ks (containing either p110 α , β , or δ) that interacts with the regulatory subunits p85, p55, and p50 (via two src **<u>2</u>** homology (SH2) domains) and specifically transduce signals of phosphorylated tyrosines from activated receptor tyrosine kinases (RTKs), and IB consisting of p110ycontaining PI3Ks that interact with the regulatory subunits p101 and p87 and are activated through interactions with the $\beta\gamma$ subunit of heterotrimeric G proteins from GPCRs (Fruman et al., 1998; Wymann and Marone, 2005). The least amount of information is known about class II enzymes, although it is believed that they may be involved in modulating and regulating clathrin distribution and assembly through formation of PtdIns-3-P, and have been found to bind PtdIns-3,4-P₂ in vitro (Wheeler and Domin, 2006). Class III enzymes exclusively produce PtdIns-3-P, which regulates vesicle trafficking by interacting with proteins containing FYVE finger domain (Fruman et al., 1998). Class I_A PI3Ks also possess protein kinase activity (in addition to the lipid kinase activity that all PI3Ks possess) and have been reported to phosphorylate many exogenous substrates such as insulin receptor substrate 1, phosphodiesterase 3B, and autoregulatory phosphorylation of Ser608 of a p85 α adaptor subunit, which serves to inhibit lipid kinase activity of the p110 catalytic subunit (Lam *et al.*, 1994; Rondinone *et al.*, 2000; Beeton *et al.*, 2000).

PI3K can be activated by cytokines and growth factors that recruit PI3K to the membrane for interaction with its phosphoinositide substrate (Cantrell, 2001; Krasilnikov, 2000). Upon formation of phosphorylated phosphoinositides, these secondary lipid messengers signal the activation of AGC (protein kinase A, G, and C) kinases such as PKB and p70S6K, which mediate responses by the regulating downstream enzymes and regulatory proteins (Brazil and Hemmings, 2001; Lizcano *et al.*, 2003). PKB has been linked to inhibition of apoptosis and differentiation corresponding with an increase in cell proliferation resulting in cancer (Franke *et al.*, 2003). Regulation of this signalling pathway is accomplished by dephosphorylation of secondary lipid messengers by <u>SH</u>2 domain-containing inositol 5'-phosphatase (SHIP) - 1/2 and PTEN (phosphatase and tensin homologue deleted on chromosome ten) phosphatases, which dephosphorylate inositides and phosphoinositides at their 5' and 3' position, respectively (Cantrell, 2001).

PI3K signal transduction is also an important component of the insulin-signalling pathway (Shepherd, 2005). The hormone insulin rapidly stimulates class I_A PI3K signalling to promote glycogen synthesis, inhibition of lipolysis, stimulation of protein biosynthetic pathways, and the translocation of the GLUT4 glucose transporter (a receptor required to allow the entry of extracellular glucose) to the plasma membrane (Shepherd, 2005; Bose *et al.*, 2002). In addition to regulating glucose metabolism and utilization, insulin signalling is also a major regulator of cell growth and development in *Drosophila* and is highly conserved from *Drosophila* to humans (Lizcano *et al.*, 2003; Bikopoulos *et al.*, 2004). *Drosophila* contains one insulin receptor (DinR) and seven insulin-like peptides, as well as homologues of mammalian insulin receptor substrates, PI3K, PTEN, PKB/Akt, and p70SK6. Because of these similarities, *Drosophila* is a useful model in the study of this pathway and its role in the coordination of cellular growth, proliferation and cell survival, with development (Bikopoulos *et al.*, 2004). Expression of *Drosophila* insulin-like peptide 2 (DILP2) results in an increase in body mass and growth in the developing eye when exclusively overexpressed in this tissue, whereas inhibition of insulin signalling corresponds to a decrease in the overall size of the fly and a reduction in the growth of imaginal discs (Edgar, 1999; Brogiolo *et al.*, 2001). These results reflect the ability of insulin signalling to inhibit the apoptosis, thereby allowing continued growth of cells and tissues by preventing cell death (Bertrand *et al.*, 1998; Lee-Kwon *et al.*, 1998; Yenush *et al.*, 1998).

It was recently determined that caffeine could inhibit the lipid kinase activity of class I PI3Ks *in vitro* at an IC₅₀ of 75 μ M for p110 δ , 400 μ M for p110 α and p110 β , and 1 mM for p110 γ . Caffeine can also block the ability of insulin to stimulate PKB with IC₅₀ values similar to those required for inhibition of PI3K activity and insulin-stimulated glucose transport in these cells (Foukas *et al.*, 2002). This suggests that caffeine could mediate its effects by directly inhibiting PI3K and its resultant downstream signalling, an effect that could also be related to caffeine's ability to induce apoptosis as inhibition of PI3K signalling could prevent PKB from inhibiting apoptosis.

Section 1-7: Caffeine and cell cycle control

Another interesting effect of caffeine is its ability to sensitize cells to radiation and various other DNA damaging or checkpoint-inducing agents. Checkpoints are regulatory mechanisms that temporarily halt the cell cycle to allow for the completion of critical events such as DNA replication, repair, or chromosomal segregation (Weinert and Hartwell, 1989). Checkpoints are crucial for ensuring that the cell undergoes an orderly, unidirectional progression through the cell cycle, ensuring that subsequent steps are completed prior to cellular division. Failure to do so results in loss of genomic fidelity (as unrepaired mutations are carried on to the next generation) and compromises genomic integrity and chromosomal stability as DNA damage caused by incomplete replication and double stranded breaks, ultimately leads to loss of chromosomal information. Observations made in a variety of cell lines treated with ionizing radiation or DNA damaging drugs revealed that checkpoint-induced delays that normally occur upon exposure to these agents, are disrupted by caffeine (Rauth, 1967; Lau and Pardee, 1982; Schlegel and Pardee, 1986). The presumed mechanism of caffeine's ability to override checkpoint responses comes from evidence suggesting that caffeine inhibits the protein kinase activity of Ataxia Telangiectasia-Mutated (ATM) and ATM and Rad3 related (ATR) kinases (Blasina et al., 1999; Sarkaria et al., 1999; Zhou et al., 2000). These kinases have been shown through genetic and biochemical evidence to be major regulators of DNA damage-induced cell cycle checkpoint and repair pathways that become activated upon detection of DNA damage or incomplete replication (Abraham, 2001). Once activated, these kinases phosphorylate downstream checkpoint effector kinases Chk1 (on S345 and S317 by ATR) and Chk2 (on T68 by ATM), which in turn phosphorylate Cdc25 on S216, as well as the inhibitory kinases Wee1 and Myt1 (Figure 1-7) (Abraham, 2001). Cdc25 is the phosphatase responsible for removing the inhibitory phosphate residues on tyrosine (Y) 15 (mediated by both Wee1 and Myt1) and T14 (mediated by Myt1) of the master regulator of mitosis Cyclin Dependent Kinase 1 (CDK1). CDK1, a complex of Cdc2 and Cyclin B, is directly responsible for starting the initial progression into mitosis by phosphorylating critical substrates, such as histones to cause condensation of chromatin in early prophase. Phosphorylation of Cdc25 at S216 prevents activation of CDK1 because it results in an allosteric movement of Cdc25 that reveals a binding site for 14-3-3 proteins and results in sequestration of Cdc25 in the cytoplasm, keeping it away from nuclear CDK1. In contrast, phosphorylation of Myt1 and Wee1 kinases by Chk1 activates these kinases to mediate inhibitory phosphorylation of CDK1. Thus, the mechanism of caffeine's ability to radiosensitize or chemosensitize cells treated with genotoxic agents is believed to be through the inhibition of ATM and ATR kinases, such that checkpoints which would normally be activated through exposure to these agents, are abrogated.



Figure 1-7. General mechanism of checkpoint control. Upon detection of DNA damage, the master regulatory kinases Ataxia Telangiectasiamutated (ATM) and ATM and Rad3 related (ATR) kinases activate downstream kinases Chk1 and Chk2, which initiate a G_2 checkpoint by inhibiting Cdc25 and the consequent activation of CDK1, which commits the drive towards mitosis. Chk1 and Chk2 are also involved in activation of inhibitory kinases Wee1 and Myt1, which maintain the checkpoint through inhibitory phosphorylation of CDK1. Caffeine inhibits the actions of ATM and ATR, resulting in the abrogation of the checkpoint and consequent premature entry into mitosis.

Section 1-8: *huc*95^{DE} was isolated in a screen for novel cell cycle checkpoint mutants

Since caffeine has the ability to override checkpoint responses by the inhibition of ATM and ATR, it has become a useful tool in the study of cell cycle checkpoint and DNA damage response pathways (Tenzer and Pruschy, 2003; Blasina et al., 1999; Sarkaria et al., 1999). This ability also made caffeine an attractive chemical to use for the possible discovery of novel cell cycle mutants, particularly if some of these mutants remained hidden due to functional redundancies within the cell cycle regulatory machinery (Silva, 2002). Control of the cell cycle is of critical importance because if left unregulated, it could lead to cancer or cell death due to genomic and chromosomal instability. Consequently, redundant mechanisms exist that can compensate for defects in components of the pathway (Abrahams, 2001). Previously, a screen was designed to discover possible novel cell cycle checkpoint mutants that may have not been detected in other screens due to possible redundancy in cell cycle checkpoint and repair systems (Silva, 2002). A precedent for this idea was suggested by the discovery of the *mus*304 gene. This gene, which was isolated from a collection of **mu**tagen sensitive Drosophila mutants, encodes a novel DNA checkpoint gene, suggesting that perhaps other regulators of the cell cycle may also be currently undetected, particularly if their functions were masked by redundant checkpoint regulation mechanisms (Henderson, 1999; Brodsky, 2000; Silva, 2002). A strategy was devised whereby the progeny of EMS-mutagenized Drosophila were assayed for the ability to develop normal eyes when they were raised in the presence of hydroxyurea (a drug that inhibits ribonucleoside reductase: an enzyme involved in the synthesis of deoxyribonucleotides) and caffeine. The rationale of the strategy was that an S-phase checkpoint would be induced by hydroxyurea (by affecting the cell's ability to replenish the depleting deoxyribonucleotide pools during DNA replication), but inhibition of the master cell cycle and DNA damage/repair pathway regulators ATM and ATR by caffeine might partially override this checkpoint response, forcing cells to undergo inappropriate mitosis unless another pathway independent of ATM and ATR could be activated to maintain the checkpoint. As a result, mutations affecting genes involved in such a redundant pathway that would normally be masked by the effects of DNA repair pathways could be uncovered. Mutations affecting genes involved in such a checkpoint response would be expected to render mutants more sensitive to these drugs, resulting in developmental defects.

Since the original premise of the screen was to discover novel cell cycle mutants, it was necessary to have a method that would allow for both the detection of such mutants and permit these mutants to be viable and fertile. Because components that regulate cell cycle checkpoint responses would target the same mitotic machinery required for all cell divisions, mutations affecting these components could often be expected to result in non-conditional lethality or sterility. To address this problem, a method that allows for creation of an adult eye composed entirely of homozygous mutant ommatidial cells in an otherwise heterozygous fly was used (Stowers and Schwarz, 1999). This method takes advantage of the GAL4-UAS system and FLP-FRT system adopted from yeast and involves driving GAL4 expression from an *eyeless* promoter (ensuring that GAL4 expression is limited to the developing eye), which would then bind to Upstream Activator Sequences fused to a FLP recombinase transgene (Golic and Lindquist, 1989; Brand and Perrimon, 1993; Stowers and Schwarz, 1999). FLP recombinase then induces site-specific mitotic recombination at FRT sites on the chromosomes within the cells of

the developing eye, resulting in twin clones of homozygous cells. Selection against one class of homozygous ommatidial cells is then accomplished by expression of a transgene carrying <u>head involution defective</u> (a gene that produces a pro-apoptotic protein) under control of a late-expressed eye-specific promoter **Glass Multiple Reporter** (GMR-hid). As a result, the development of any cell inheriting *GMR-hid* is terminated, leaving only cells that are homozygous for the mutagenized chromosome (see Figure 1-8). The Drosophila compound eye is an ideal organ for such a screen, as it is not required for viability of the organism and is an easily viewed structure, allowing many different mutant phenotypes affecting size, texture, and organization to be distinguished. The eye also has a well-established (albeit complex) system of development, making it an excellent organ to study how pattern formation is regulated by the control and timing of cell proliferation and differentiation. The adult eye is composed of approximately 700 repeating facets or ommatidia, each of which is generated from 19 precursor cells that develop within the eye-antennal imaginal disc (Wolff and Ready, 1993). Each ommatidium, responsible for focusing and transmitting light from a small portion of the visual field to the optic regions of the brain, consists of a precise concentric arrangement of 8 photoreceptor cells surrounded by accessory cone and pigment cells. From embryogenesis until the end of the second larval instar, cells in the eye-antennal disc proliferate in an unpatterned manner. During the last larval instar, ommatidial specification and retinal differentiation becomes evident through the appearance of a dorsal-ventral invagination called the morphogenetic furrow that migrates posteriorly to anteriorly along the apical surface of the eye-antennal disc (Figure 1-9). Cellular proliferation in the anterior part of the eye imaginal disc still remains mitotically

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Figure 1-8. Method for generating homozygous ommatidial mitotic clones using the EGUF-GMR-hid system. GAL4 expression is driven from the *eyeless* promoter, restricting expression of GAL4 to ommatidial cells, where it interacts with upstream activating sequence (UAS) to drive expression of a Flp recombinase. Flp recombinase mediates homologous recombination at FRT sites and expression of the dominant death gene *hid*, driven by the Glass Multiple Reporter (GMR) promoter to restrict expression of this gene within the developing eye, ensures that only cells inheriting homozygous combinations of the chromosome of interest (designated by the chromosome arm with an asterisk) survive to undergo the next round of replication. After several generations, the result would be a *Drosophila* eye composed entirely of homozygous clones for this chromosome. Results from both an odd and even number of recombination events are depicted, along with the predicted percent chance of these cells producing homozygous or heterozygous progeny cells (from Stowers and Schwarz, 1999).



Figure 1-9. Development of the *Drosophila* eye imaginal disc. During the first and second larval instars, cells within the developing eye are mainly proliferative and governed by individual cell cycles (blue dots indicative of cells in S-phase and red dots indicative of cells undergoing mitosis). During the last larval instar, the morphogenetic furrow (MF – green bar) designating retinal differentiation and specification migrates in a posterior to anterior direction, synchronizing and arresting cells in G₁ that are caught in its wake. Within the furrow, ommatidial precursors are specified and organized into preclusters for differentiation into ommatidia, while cells that are not organized undergo a single round of mitosis forming a second mitotic wave prior to differentiating (note: proliferative cells ahead of the furrow are often referred to as the first mitotic wave even though these cells have no obvious synchrony or pattern).
unpatterned and uncoordinated. Directly ahead of the furrow however, cells proceed through S-phase then undergo a semi-synchronized mitosis referred to as the first mitotic wave (FMW) (Wolff and Ready, 1993). Upon exit from mitosis, these cells become arrested in G_1 as they enter the furrow. Within the furrow, the G_1 arrested ommatidial precursors become specified and organized into five-cell pre-clusters that will eventually differentiate into an adult ommatidium (Wolff and Ready, 1993; Baker, 2001). Behind the morphogenetic furrow, the cells of the pre-clusters undergo neuronal differentiation and never divide again while the remaining unspecified cells surrounding the G₁-arrested ommatidial pre-cluster cells synchronously undergo a round of S-phase and mitosis referred to as the second mitotic wave (SMW). These cells will then be recruited to make up the remaining 14 precursor cells required for a complete ommatidium. The SMW is necessary to generate enough precursor cells to complete each ommatidium. Supernumerary cells generated at this stage are destroyed later in the pupa by apoptosis, ensuring that a precise number of cells are maintained for development of the adult eye (Wolff and Ready, 1993; Baker, 2001). Because the eye is characterized by specific requirements and a temporal pattern of cell cycle regulation, it is an amenable system for identifying mutations causing defects in cell cycle regulation.

Using the method described earlier, referred to herein as the EGUF-GMR*hid* method for <u>eyeless-G</u>AL4; <u>UAS-F</u>LP, 3 of the 5 large chromosome arms in *Drosophila* (2L, 2R, and 3R, respectively) were screened (see Figure 1-10 for an example of the screening procedure for 2L). From the initial cross, F_1 males that displayed a rough-eye phenotype in the presence of 3 mM hydroxyurea (HU) and 2 mM caffeine were re-tested on normal media to identify conditional mutants, which were then confirmed and mapped

further. From this screen, a total of 9 mutations (herein referred to as *huc* mutants for *hydroxyurea caffeine-dependent* mutants) were isolated, with 8 mutations mapping to the left arm of the second chromosome (that were subsequently separated into 5 complementation groups), and 1 mutation mapping to the right arm of the third chromosome. This 3R single mutant was mapped by deficiency mapping and complementation analysis to cytological region 95D-E (and was thus referred to as *huc*95^{DE}). The *huc*95^{DE} mutant displayed a severely reduced and disorganized eye with irregular and fused ommatidia when larvae were grown on media containing HU and



Figure 1-10. A genetic screen of chromosome 2L to identify mutants sensitive to hydroxyurea and caffeine during eye development. Eye-specific recombination between centromerically located FRT sites, in combination with eye-specific expression of the apoptotic inducing gene *hid*, results in eyes that are essentially a clone of a single genotype; the genotype of the remainder of the organism is heterozygous. F_1 larvae are grown in the presence of HU and caffeine. Males exhibiting an eye phenotype are selected and crossed again to EGUF females and progeny are grown in the absence of HU and caffeine. Male progeny from this cross that exhibit a wild-type or near wild-type phenotype were isolated for further analysis. Similar methods were applied in screening chromosomes 2R and 3R, using arm-specific FRT stocks (from Silva, 2002).

caffeine. This phenotype was more severe than any of the other mutants mapped on the second chromosome (see Figure 1-11). It was later discovered that treatment of $huc95^{DE}$ with caffeine alone produced this severe eye phenotype, whereas treatment with hydroxyurea alone produced eyes that were similar to wildtype, suggesting that $huc95^{DE}$ are primarily sensitive to caffeine. When $huc95^{DE}$ hemizygous larvae are viable were reared on standard molasses media. $huc95^{DE}$ hemizygotes however, displayed caffeine-dependent lethality when grown in media containing caffeine concentrations as low as 0.5 mM, suggesting that the gene associated with this mutation was not essential, but was conditionally required for viability in the presence of caffeine.

To determine if these defects occurred as a result of cell cycle abnormalities during early stages of ommatidial differentiation, imaginal eye discs were dissected from EGUF/+; FRT82B huc95^{DE}/FRT82B GMR-hid3R third instar larvae and analyzed using antibodies to a neuronal RNA-binding protein called <u>embryonic lethal</u>, <u>abnormal visual</u> (ELAV), which stains all differentiated neuronal cells (Silva, 2002; Yao *et al.*, 1993). Imaginal discs from trans-heterozygous mutant larvae grown in the presence of caffeine were smaller and more fragile than heterozygous controls and ELAV staining revealed that differentiation of the ommatidial preclusters still occurs with no change in size of the individual pre-clusters of trans-heterozygous mutant eye discs (although the pattern appeared to be slightly disrupted) (Figure 1-12). This suggests that the decreased size of the eye imaginal discs is due to fewer cells being present as opposed to smaller cells within the disc. The eye imaginal discs were also treated with Bromodeoxyuridine (BrdU), a base analogue of thymidine that is incorporated during DNA synthesis, identifying cells undergoing S phase. Third instar larval eye imaginal discs from trans-



Figure 1-11. Scanning electron micrographs of adult eyes from individuals of the genotype EGUF/+; FRT82B huc95E-D/ FRT82B GMR-hid3R CL3R grown in the absence (A) and presence (B) of caffeine. The phenotype depicted here is indistinguishable from the phenotype that is observed when individuals are grown in the presence of both HU and caffeine. Eye imaginal discs from individuals of the genotypes EGUF/+; FRT82B huc95E-D/ TM3 Ser (C and E) and EGUF/+; FRT82B huc95E-D/ FRT82B GMR-hid3R CL3R (D and F) grown in the presence of caffeine. In the presence of caffeine, eyes are severely reduced in size and ommatidia are fused and greatly reduced in number (compare A and B). Staining with elav antibody reveals neuronal differentiation still occurs. Discs are smaller at third instar compared to heterozygous controls (compare C and D) but individual cells are not reduced in size (compare E and F) (from Silva, 2002).

heterozygous *huc*95^{DE} mutants and heterozygous controls treated with caffeine and HU fewer cells stained with BrdU in the presence of these drugs were found to be comparable to controls (see Figure 1-12) (Silva, 2002). In addition, the pattern of staining ahead and behind of the morphogenetic furrow seemed to be disrupted and more diffuse in the presence of caffeine, suggesting that this mutant may have cell proliferation defects during ommatidial development, or exhibit aberrant proliferation due to an increase in apoptotic responses.

Mapping of $huc95^{DE}$ by E. Silva identified 25 potential candidates within this region, which was reduced to 22 as a result of complementation tests with mutants of three genes: *Syntaxin 1A* (*Syx1A*), *Adenomatous polypopsis coli 2* (*Apc2*), and *Tuberous sclerosis complex 1* (*Tsc1*). Because of the severity of the $huc95^{DE}$ phenotype, and the fact that this mutant showed defects that could be due to effects on cell cycle regulation, the gene associated with this mutant became an attractive candidate to pursue. As a result, my project was developed to identify the gene associated with the $huc95^{DE}$ mutation, determine the nature of the mutation, and carry out further phenotypic analysis to elucidate the specific defects caused by the mutation.



Figure 1-12. Eye imaginal discs from individuals of the genotypes EGUF/+; FRT82B huc95E-D/ TM3 Ser (A and C) and EGUF/+; FRT82B huc95E-D/ FRT82B GMR-hid3R CL3R (B and D) grown in the presence of caffeine and labeled with the nucleoside analog BrdU (from Silva, 2002).

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Chapter 2: The Mapping and Identification of huc95^E

Section 2-1: Introduction

In order to elucidate the molecular nature of the $huc95^{DE}$ mutation and to determine the reasons for caffeine sensitivity, it was necessary to identify which gene on the right arm of the third chromosome was associated with this phenotype. Furthermore, determining the identity of $huc95^{DE}$ could help to assess the success of this screen regarding its original intent, which was to identify novel cell cycle checkpoint functions. Originally, *huc*95^{DE} was mapped cytologically by deletion mapping and complementation analysis to the cytological region of 95D11-95F15, because $huc95^{DE}$ failed to complement the deficiencies Df(3R)crb87-4 (extending from 95D11-E2 to 96A2) and Df(3R)crbF89-4 (extending from 95D7-11 to 95F15), respectively (Silva, 2002). Further refinement of the position to 95D11 to 95E6-7 was determined by complementation testing of $huc95^{DE}$ with the synthetic deficiency Df(3R)slo3, which has a large deletion on the right arm of the third chromosome (extending from 94D4-10 to 96A18) and the proximal portion (94D4-10 to 95E7-F1) of this deleted region duplicated, inverted, and inserted into the second chromosome at position 57B3-5 (see Figure 2-6A). These results had narrowed the number of potential candidates down to 26 genes (see Table 2-2). Out of this initial list, huc95^{DE} was found to complement the genes Svntaxin 1A (Svx1A at position 95E1), Tuberous sclerosis complex 1 (Tsc1 at 95E4), and Adenomatous polyposis coli 2 (Apc2 at 95E6), thereby limiting the number of candidates at the start of this project to 23 predicted genes within this region.

Besides deficiency mapping and complementation analysis, several other strategies are available to further narrow down and map *Drosophila* mutations in a cost

effective and efficient manner. One such strategy is the genetic mapping of the relative positions of genes by recombination frequency analysis. This can be done with a classical three point cross whereby the position of a mutation is mapped with respect to two other mutations on the same chromosome. Another method that is useful in mapping is to mobilize P-elements to synthesize flanking deletions that when mapped, can be used to narrow down the location of a gene expressing a particular trait. This method involves isolating P-element-induced recombinants and using segregation patterns of recessive markers flanking the mutation to determine whether the mutation segregates proximal or distal to the site of insertion (Chen et al., 1998). Determination of the molecular breakpoints of flanking deletions of recombinants that segregate with the mutation can then be used to narrow down the region to where the mutation exists on the chromosome. This is an informative method, as even determining the breakpoints of recombinants that do not segregate with the mutation can also be used to exclude candidate genes lying within the region. Another method is the molecular analysis and sequencing of candidate genes. This method takes advantage of the completed Drosophila genome sequence of predicted genes and allows for the identification and analysis of sequence alterations in a search for putative candidates. This method could not only identify huc95^{DE}, but also elucidate the molecular nature of the mutation associated with the gene. Although it did not seem to be very practical to sequence all 23 candidates within this initial region, this method would become useful if this region was further narrowed down by any of the other methods previously described (so that only a few remaining candidates would remain to be sequenced), or could become useful in the confirmation and analysis of a particular candidate. Furthermore, obvious candidates for which a plausible mechanism

could be proposed for caffeine sensitivity (such as gene candidates involved in apoptosis, caffeine metabolism, or genes involved with cellular proliferation or cell cycle checkpoints) could be directly analyzed by this method in the absence of any available mutants for these genes. Each of the above methods were attempted in order to determine the location of the *huc*95^{DE} mutation. This chapter will discuss each approach and the information that I obtained from them.

Section 2-2: Materials and Methods

Section 2-2a: Fly stocks and crosses

All *Drosophila* mutants and fly stocks used in the experiments described within this chapter are listed in Table 2-3. All fly stocks and crosses (unless otherwise stated) were raised on standard molasses and cornmeal media and grown either at room temperature or at 25°C in either vials or plastic fly bottle with volumes of 6 ml and 60 ml media, respectively. For crosses involving caffeine treatment in vials, caffeinecontaining food was prepared by scoring the media with a clean metal spatula and adding 250 μ L of caffeine (dissolved in distilled water) at an appropriate concentration (normally from a 50 mM caffeine stock solution for a final concentration of 2 mM) to yield the desired concentration in vials. This volume was previously empirically calculated to ensure complete absorption of the caffeine. Prepared vials were allowed to soak over night after which flies were tipped into this media and allowed to lay eggs for approximately 24 hours prior to tipping again to a new vial. Alternatively, fly crosses were set up within the vials and allowed to lay eggs for 24 hours after which time the flies were tipped into a new vial. The media was then scored and caffeine added as described above. This method was done to address any concerns about flies not laying in caffeine-tainted media because of perceived sensitivity to the media. However, both methods yielded similar results and approximately equivalent amounts of progeny, indicating that either method was suitable for the caffeine experiments described in this chapter.

*huc*95^{DE} hemizygotes were originally produced by either crossing *huc*95^{DE}/*TM3 Ser-GFP* to either *Df*(3R)*crb87-4*/*TM3 Ser-GFP* or *Df*(3R)*crbF89-4*/*TM3 Ser-GFP*, and selecting non-Ser adults or non-GFP third instar larvae. Because of the incomplete penetrance and cryptic nature of the *Ser* marker, and problems involving damage or loss of wings in the media, crosses involving *huc*95^{DE}/*TM3 Sb*, *Ser* and *Df*(3R)*crb87-4*/*TM3 Sb*, *Ser*, *Df*(3R)*crbF89-4*/*TM3 Sb*, *Ser*, or *Df*(3R)*W6*/*TM6C Sb*, *Tb* were later used, selecting for absence of Sb in adult flies. To detect hemizygous 2nd and 3rd instar larvae, crosses involving *huc*95^{DE}/*TM6B Sb*, *Tb*, *e* and *Df*(3R)*crb87-4*/*TM6B Sb*, *Tb*, *e* were used, selecting for non-Tb larvae. Crosses to generate deletion mutants through *P*element mobilization are outlined in Figure 2-3.

Section 2-2b: PCR and sequencing

DNA used in PCR and sequencing reactions was obtained by homogenizing adult flies of the appropriate genotype in genomic DNA homogenization buffer (100 mM NaCl, 0.2 M sucrose, 100 mM Tris pH 9.5, 50 mM EDTA, and 0.05% SDS) and incubated at 65° C for 30 minutes, followed by addition of 8 M of potassium acetate and further incubation on ice for 30 minutes. Following centrifugation at ~ 12000 rpm (17211 xg) for 10 minutes, 0.7 volumes of isopropanol was added to the supernatant, incubated on ice for an additional 30 minutes, and centrifuged at ~ 8000 rpm (7649 xg) for 10 minutes. The pelleted DNA was then washed in 70% ethanol, resuspended in TE Buffer (10 mM Tris-Cl pH 7.5 and 1mM EDTA), and treated with RNase A at 37°C for 30 minutes. RNase and any remaining protein were removed through phenol-chloroform extractions (as described in Sambrook et al., 1989), followed by isopropanol/ethanol precipitations (as described above but using 7.5 M ammonium acetate), and resuspension in sterile water. The resultant DNA was then quantified by spectrophotometry or gel quantification analysis. All PCR reactions were carried out for 30 cycles utilizing the appropriate primer pairs for given genes and sequences at concentrations of 1 mM (see Appendix for list of primers). PCR reaction mixes containing 10X PCR Buffer, 50 mM MgCl₂ solution, and *Thermus aquaticus* DNA polymerase were obtained from Invitrogen, as well as all primers used. Sequencing reactions were performed using the Big Dye Terminator V3.1 sequencing mix and sequencing primers listed in the appendix for the various genes listed. For large genomic regions, overlapping template sequences of approximately 2000 to 3000 nucleotides were amplified. Sequencing reactions were conducted using 100 - 500 ng of template DNA, 5 pmol of sequencing primer, 4 μ L of sequencing pre-mix buffer (consisting of 200 mM Tris, pH 9.0 and 5 mM MgCl₂), and 4 mL of the BigDye premix in a total volume of 20 µL. PCR reactions were then run with a denaturation temperature of 95°C for 30 sec, annealing temperature of 50°C for 15 sec, and an extension temperature of 1 min for 30 cycles. The Molecular Biology Sciences Unit at the University of Alberta then resolved the sequencing reactions by gel electrophoresis using an ABI 3700 automated DNA sequencer, and the resulting sequences were analyzed using the sequence analysis program Gene Tools 2.

Section 2-2c: Embryo DNA Preparation and PCR

The extent of the Df(3R)W6 deletion was determined using DNA isolated from embryos collected on grape juice plates after 1 day of egg laying from a Df(3R)W6/TM3CSb, Tb stock. Each embryo was then macerated in 10 µL of homogenization buffer (10 mM Tris-HCl pH 8.0, 1 mM EDTA pH 8.0, 25 mM NaCl, and freshly added Proteinase K at 200 µg/ml), incubated for 30 minutes at 37°C (for Proteinase K digestion), followed by heat inactivation of Proteinase K at 99°C for 10 minutes. Df(3R)W6 homozygote embryo DNA samples were identified by the inability to amplify fragments from Apc2and Tsc1, known to reside within the deficiency, and the ability to amplify fragments from Myt1, which was known to reside outside of the deleted region. 1 µL of this mixture was immediately used as a template in PCR reactions performed as described above utilizing the primer sets for Myt1, Tsc1, Apc2, mRpS24, CG13605, CG13606, CG13607, CG6129, CG6178, CG6182, CG5463, CG5510, CG18428, and CHORD (see appendix).

Section 2-2d: RT-PCR and RNA isolation

RNA for RT-PCR analysis was isolated using the Qiagen RNAasy Mini Kit (catalogue number 74104). Total RNA was obtained by macerating adult flies or larvae of the appropriate genotype in the provided lysis buffer using a mortar and pestle, and homogenizing the lysate by passage through a 20 gauge sterilized needle. The resultant suspension was then processed directly according to the protocol provided with the kit for isolation of total RNA. The isolated RNA was then used for RT-PCR using the One Step RT-PCR Kit (catalogue number 210210) and subjected to reverse transcription reaction for 30 minutes at 50°C, an initial PCR activation step of 95°C for 15 minutes, followed by

PCR denaturation, annealing, and extension reactions at appropriate temperatures as described above for 40 cycles. A final extension time of 10 minutes at 72°C was also programmed to ensure complete extension of desired templates. Results were then analyzed by gel electrophoresis on a 0.5% Tris Borate EDTA (TBE) or 1% Tris Acetate EDTA (TAE) agarose gel.

Section 2-2e: Synthesis and mapping of transgenic rescue lines

The restriction enzymes EcoRI, ScaI, EcoRV, and XhoI, and T4 DNA ligase, all used in the cloning of a myo95E rescue construct, were obtained from Invitrogen. Genomic DNA for the myo95E genomic rescue construct was excised from the Bacterial Artificial Chromosome BACR29F06, obtained from BACPAC resources at the Children's Hospital Oakland Research Institute. The genomic rescue plasmid for myo95E was constructed by excising a 8039 bp fragment from BACR29F06 using a blunt ended restriction enzyme Scal, and a sticky ended cutter XhoI (Figure 2-12). This fragment was predicted by the *Drosophila* genome project to encompass the whole myo95E gene (including all 15 introns, 3', and 5' untranslated regions) as well as most of the downstream region up to the next gene (but did not contain the upstream genomic region). To add an EcoRI restriction site, which would be used for insertion into the transformation vector pUAST, this 8039 bp fragment was subcloned into the vector pBluescript cut with EcoRV and XhoI. To include regulatory elements upstream of myo95E, a 3672 bp fragment containing the upstream genomic region was also excised from BACR29F06 using XhoI restriction sites, and ligated to the pBluescript plasmid containing the 8039 bp fragment restricted with XhoI. Plasmid contructs containing the 3672 bp fragment ligated within the correct orientation were identified by PCR using

primers that flanked the ligation site, and the resulting 11711 bp fragment (8039 bp + 3672 bp) was sequenced to confirm its integrity. The 11711 bp fragment was then excised using EcoRI and XhoI, and cloned into the transformation vector pUAST (described by Brand and Perrimon, 1993). Transformation through injection into preblastoderm embryos was carried out as described by Rubin and Spradling (1982) with the construct concentrated and purified to a concentration of 5 μ g/10 μ L, along with a Δ 2-3 helper plasmid at 1 μ g/10 μ L that was microinjected into stage 4 bleach-dechorionated embryos from yw flies. First instar survivors were then raised on instant Drosophila media, and were subsequently crossed to yw flies as adults. Two putative transformants (identified by orange eye colour due to expression of the white gene present in the Ptransformation vector pUAST) were obtained and the rescue construct was subsequently mapped in both transformants to the third chromosome. Because *huc*95^{DE} also mapped to the third chromosome, recombinants were generated using the strategy shown in Figure 2-13A that contained the genomic rescue construct and Df(3R)crb87-4 on the same chromosome. Putative recombinants were selected on the basis of inheritance of $P[w^+]$ transgene carrying the rescue construct and e from the deficiency chromosome with a third chromosomal balancer and then tested for complementation against the deficiency to identify recombination events that linked the construct (evident by the presence of $[w^{\dagger}]$ gene expression) and the *e* recessive mutation without the deficiency. Recombinant chromosomes carrying both the transgene and deficiency were then tested for complementation with *huc*95^{DE} mutants in the presence and absence of 2 mM caffeine, as described in Figure 2-13B.

Section 2-3: Results

Section 2-3a: Mapping of huc95^{DE} through Recombination

Because of the caffeine-sensitive conditional lethality of *huc*95^{DE} hemizygotes, a strategy was devised to map $huc95^{DE}$ through the classical method of constructing three factor crosses and determining recombination frequencies. This strategy involved the use of two unconditional recessive embryonic lethal mutants for the genes syntaxin 1A (syx1A) and crumbs (crb) (mapping to 95E1 and 95F10-11, respectively), as well as the deficiency Df(3R)crb87-4 which spanned the region containing these genes and $huc95^{DE}$ (see Figure 2-1 for the crossing scheme). Syx1A is a presynaptic membrane protein present in several tissues but most highly expressed within cells of the nervous system where it is thought to act as a receptor for synaptic vesicle docking and fusion (Broadie et al., 1995). syx1A null mutant embryos fail to secrete most or all cuticle, exhibit abnormal gut morphology, and have subtle defects in axonal morphology (with axon bundles of segmental and intersegmental nerves appearing thickened and irregular in shape). In addition, there is a complete failure to evoke endogenous neurotransmitter release, resulting in an absence of synaptic transmission (Schulze et al., 1995). The crb gene encodes an apical transmembrane protein that is involved in determining the apicalbasolateral polarity of epithelial cells - an important process in the organization and establishment of compartmentalization in multicellular organisms (Tepass et al., 1990). Consequently, crb mutants exhibit embryonic lethality shortly after gastrulation (a period when extensive epithelial movement and reorganization occurs to form the three germ layers, eventually giving rise to body axis and organogenesis).

The fact that both syxIA and *crb* mutants were able to complement $huc95^{DE}$ allowed for a strategy to be devised whereby the caffeine sensitivity phenotype could be used to map $huc95^{DE}$ with respect to these two genes (see Figure 2-1). In this strategy, the frequency of forming huc95^{DE}-syx1A, huc95^{DE}-crb, and syx1A-crb recombinant double mutants was determined by crossing flies with the genotypes huc95^{DE}/syx1A, huc95^{DE}/crb, and syx1A/crb to Df(3R)crb87-4/TM3 Sb, Ser in the presence of caffeine and looking for progeny that did not inherit the TM3 balancer. Since the huc95^{DE} hemizygotes exhibited caffeine-sensitive lethality while *crb* and *syx1A* hemizygotes exhibit embryonic lethality, the only non-balancer progeny that would survive would be those that have previously undergone a recombination event to create a wildtype chromosome (Figure 2-1). Using a concentration of 2 mM caffeine (the concentration used in the screen that resulted in the isolation of these mutants), crosses involving *huc*95^{DE}/ *crb* females with *Df*(3R)*crb*87-4/ TM3 Sb males resulted in a recombination frequency of 0.021 (n=5007) for flies not containing the Sb balancer. Crosses between huc95^{DE}/ syx 1A females with Df(3R)crb87-4/TM3 Sb males resulted in a recombination frequency of 0.07 (n=3937), while crosses between syx 1A/ crb resulted in a recombination frequency of 0.017 (n=9541). Analysis of these frequencies suggested that huc95^{DE} might be closer to the crb locus than the syx1A locus (0.021 cM from $huc95^{DE}$ to crb as compared to 0.07 cM from $huc95^{DE}$ to syx 1A). However, the map distance determined between the loci of syx 1A and $huc95^{DE}$ (0.07 cM) did not add up to the sum of the proposed distances of svx 1A to crb, and crb to huc95^{DE} (0.038 cM) (Figure 2-2) suggesting possible inaccuracies in the data. Furthermore, these results suggested that *huc*95^{DE} mapped to the right of *crb* (which is located cytologically at 95F10-11), which conflicts with the original mapping of huc95^{DE} to the cytological regions of 95D-E through deficiency mapping. An explanation for these anomolous results is that perhaps some of the recombinants isolated were $huc95^{DE}$ hemizygotes who escaped the lethal effects of caffeine and managed to complete development to adulthood. This would artificially boost the recombination frequency by increasing the number of putative recombinants, thereby increasing the proposed map distance between the respective genes. To test for this, male recombinants from each of these crosses were collected and subjected to single male crosses with Df(3R)crb87-4/TM3 Sb virgin females in the presence and absence of 2 mM caffeine to test if they were true recombinants. If these flies were recombinants, then the non-balancer progeny of such a cross should be able to grow and develop normally on caffeine-containing medium. Alternatively, if the flies were escapers, non-balancers would be susceptible to caffeine sensitivity and not be sensitive in its absence. Out of 10 putative $huc95^{DE}/crb$ recombinant males tested, 5 behaved as recombinants while 5 behaved as escapers. Similar results were also obtained with crosses done with 23 huc95^{DE}-syx 1A putative recombinants, which yielded 10 possible escapers and 4 recombinants (with the remaining unclassifiable due to lack of progeny in these crosses). From these tests, it could be concluded that the genetic map is most likely affected by escapers artificially increasing the recombination frequency, which supports the inability of the map distances to add up as well as the original mapping data.

Section 2-3b: Mapping of *huc95^{DE}* through P-element mobilization and flanking deletion synthesis

Another strategy to map the location of the $huc95^{DE}$ mutation was to synthesize flanking deletions by mobilizing transposable *P*-elements within the 95D-E region. To

synthesize a library of flanking deletions, stocks carrying four different P-element insertions mapping within the cytological regions of 95D-E were crossed to males containing a transposase source and third chromosomal recessive markers claret (ca, located cytologically at 99B8-10) and ebony (e, located cytologically at 93C7-D1). The *P*-element lines used for this experiment were $P\{EP\}Syx1A^{EP3215}$ (mapped cytologically to 95E1, inserting 556 bp upstream from syx 1A), P{EP}CG6178^{EP3637} (mapped cytologically to 95E5-6, inserting 206 bp upstream of CG6178 and 820 bp upstream of Mpk2), P{EP}mask^{EP601} (mapped cytologically to 95F3-5, inserting 2995 bp within the gene <u>multiple</u> <u>ankyrin repeats</u> <u>single</u> <u>KH</u> domain (mask)), and $P{GT1}CG33100^{BG01713}$ (mapped cytologically to 95E1, inserting 3197 bp upstream of syx1A). Males containing the transposase source and one of these P-element insertions, were then crossed to homozygous *ca* and *e* females to detect *P*-element transposase-associated recombination events, scored by the absence of either *ca* or *e* in the resulting progeny (see Figure 2-3). Crossing these recombinants to *huc*95^{DE} mutants in the presence and absence of caffeine to identify recombinants that were caffeine sensitive could then identify useful recombinant lines containing flanking deletions that deleted *huc*95^{DE}. The putative deletions within these caffeine-sensitive recombinants would then be molecularly mapped by inverse PCR to narrow down the region of interest. In such a strategy, mapping the breakpoints of deletions of recombinants that do not exhibit caffeine lethality would also be informative, as mapping these would allow the exclusion of candidate genes from the region.

Males containing the transposase source on the second chromosome with the balancer CyO, and a third chromosomal dominant mutation Sb over a 2-3 translocation

dominant mutation (apterous xasta (ap^{xa})), were crossed to females from a homozygous stock consisting of e and ca markers (Figure 2-3). Males from this cross were identified by the inheritance of Sb (and thus, the transposase source as a result of co-segregation of the second and third chromosomes in this cross due to the translocation) and crossed to each of the four *P*-element lines. Resultant progeny males, which had inherited the transposase source and P-element, were crossed again to the homozygous e, ca stock to identify events of recombination that presumably would have taken place within these males. Upon examination of all progeny from several trials of this final cross (amounting to up to 5 bottles for each P-element line), no recombinants were obtained (as evidenced by the inability to recover only ca or e progeny). It was also noticed that in flies harbouring both the transposase source and the *P*-element, the expected mottling of eye colour (whereby pigmentation within the eye appears patchy due to position effect variegation) was not evident, suggesting that the transposase source used might be very weak or even non-functional. This would explain the difficulty in obtaining male recombinants since this strategy had depended upon functional transposase expression in order to induce *P*-element excision events. To address this problem, another stock with a stronger transposase transgene inserted on the third chromosome and known to transpose efficiently was obtained and used in the crossing scheme presented in Figure 2-4. In this scheme, males containing the transposase source marked with Sb were crossed to the Pelement insertion line and then crossed to a stock with a third chromosomal balancer that did not contain the dominant mutation Sb. The resulting progeny would then be crossed to the $huc95^{DE}$ stock (or a deficiency that deleted $huc95^{DE}$) and tested for caffeine sensitivity to determine if they complement $huc95^{DE}$. Such flies could carry a new deletion containing $huc95^{DE}$ caused by excision of the *P*-element. This strategy would not be as informative as the previous one, since negative results could not also be used to rule out possible gene candidates. This method did, however, have the advantage of taking fewer generations of crosses to obtain results since no recessive markers would have to be crossed in to identify recombinants. Unfortunately, all attempts to generate males containing the transposase insertion and the *P*-element line $P\{EP\}CG6178^{EP3637}$ failed, possibly due to second site lesions which might have existed on both third chromosomes harbouring the *P*-element and transposase source. Since this *P*-element corresponded to a gene that mapped within the smallest region that $huc95^{DE}$ had been mapped to (defined by Df(3R)W6, as discussed later), excision events from this *P*element line had the greatest opportunity of generating useful deletions with which to map $huc95^{DE}$. Rather than testing the other lines or devising alternative strategies for recombinational mapping, more time and focus was devoted to other types of experiments as discussed below.

Section 2-3c: Deficiency Mapping and Complementation Analysis

When I started this project, $huc95^{DE}$ was already mapped to cytologically to 95D-E by crosses to chromosomal deletions. Complementation tests with mutants affecting three genes in this region limited the candidates to 23 genes. To further narrow down the number of candidates, a complementation test was done with Df(3R)crb87-4 (a deficiency extending from 95D11-E2 to 96A2 that did not complement $huc95^{DE}$) and Df(3R)mbc-R1 (extending from 95A6-7 to 95D6-11 and did complement $huc95^{DE}$) to see if these deficiencies overlap. These two deficiencies failed to complement, suggesting that they overlap. Since Df(3R)mbc-R1 complements $huc95^{DE}$, if the extent of the overlap of this deletion could be determined, then genes within this overlap could be eliminated as candidates. syxIA, one of the previous genes which was tested and shown to complement the $huc95^{DE}$ mutation and which mapped cytologically to 95E1, was unable to complement Df(3R)mbc-RI, suggesting that this gene resided within this region. As a result, any possible candidates which localized upstream of syxIA could be excluded since this gene was localized in a deficiency which complemented $huc95^{DE}$. These results eliminated one gene upstream of syxIA, further refining the number of candidates to 22 genes.

The region where $huc95^{DE}$ maps was further narrowed down by complementation tests with another deficiency called Df(3R)W6, reported to delete the gene *Adenomatous polyposis coli 2 (Apc2)* which mapped to 95E6 (Ahmed *et al.*, 2002). $huc95^{DE}$ failed to complement Df(3R)W6, suggesting that Df(3R)W6 contained the gene associated with caffeine sensitivity. Complementation tests involving Df(3R)W6 and mutants of the *Tsc1* gene (mapped at 95E4) and the *Syx1A* gene (mapped at 95E1) had showed that Df(3R)W6failed to complement 4 different alleles of *Tsc1*, but did complement an allele of *Syx 1A*, suggesting that the deletion extended cytologically to 95E4 (deleting *Tsc1*), but not past 95E1. To further narrow down the possible candidates, the next logical step was to determine the approximate extent of the W6 deletion and the genes that it contained.

Because Df(3R)W6 was maintained over a balancer containing the larval marker Tubby (Tb) (which results in shorter third instar larvae and as a result, shorter pupal cases), and because no non-Tb pupal cases were seen in any of the stock bottles, Df(3R)W6 was lethal in a homozygous state (which was not surprising since this deficiency deleted Tscl and Tscl homozygotes die as second instar larvae). If Df(3R)W6

homozygotes were capable of surviving as embryos however, it would be possible to obtain homozygous Df(3R)W6 embryonic DNA to test for the presence or absence of specific genes by PCR amplification (Figure 2-5A). To do this, DNA was extracted from embryos deposited over a 12 hour collection period and Df(3R)W6 homozygotes were identified by two criteria: 1) the ability to amplify a 1650 bp fragment from Myt1 (localized to 64E7, as a positive control) and 2) the inability to amplify 575 bp and 540 bp fragments from Tsc1 and Apc2 since genetic tests described earlier had demonstrated that Df(3R)W6 contained these genes (Figure 2-5B). Subsequently, systematic tests were conducted to determine that Df(3R)W6 deleted the genes CG13607 and CG5510, but not CG13605 or CG13606, genes upstream and downstream from CG13607 and CG5510, respectively (see molecular evidence - Figure 2-6). Furthermore, PCR tests for CG18428 and CHORD (genes upstream and downstream of CG13605 and CG13606, respectively) had also shown amplification, confirming that Df(3R)W6 did not extend past CG13605 and CG13606 (data not shown). Even though both CG13605 and CG13606 displayed amplification, the possibility still existed that Df(3R)W6 could have extended into the 5' untranslated region or important upstream regions that were important for expression and function of these genes. As a result, these genes were still considered as candidates and defined Df(3R)W6 as extending cytologically from 95E1 to 95E7-E8, a region containing 18 predicted genes (see Figure 2-7B). Since Apc2 and Tsc1 were previously shown to complement *huc*95^{DE} mutants (see Table 2-2), the number of potential candidate genes for the *huc*95^{DE} mutation was reduced to 16 genes from a total of 23 original candidates, which prompted a renaming of the $huc95^{DE}$ mutant to $huc95^{E}$.

For further analysis of candidate genes within the region, P-element insertion lines associated with the genes kal-1, CG6164, sec10, CG6178, and CG6182 were obtained and tested against $huc95^{E}$ for complementation. The gene kal-l is the Drosophila homologue of the extracellular matrix cell-adhesion protein anosmin-1, which is associated with the human X-linked genetic disorder Kallmann syndrome (Andrenacci et al., 2004). This disease is characterized by anosmia (no sense of smell), hypogonadism, and sterility, and is thought to arise because anosmin-1 is suspected to play a role in the formation of the olfactory bulb and in the targeting and migration of axons of olfactory neurons and gonadotropin-releasing-hormone neurons into the brain (thus explaining the phenotypes as gonadotropin is the hormone responsible for gonad and genital development) (Lutz et al., 1993; Hardelin et al., 1999). Sec10 is a component of the exocyst complex, which is an important macromolecule in the secretory pathway, purported to play a role in the trafficking of secretory vesicles to the plasma membrane (Lloyd et al., 2000). The genes CG6164, CG6178, and CG6182 are novel genes with no known function or homologues that are also $huc95^{E}$ candidates because they reside in the region deleted by Df(3R)W6. The insertion points for the *P*-insertion elements associated with kal-1, CG6164, sec10, and CG6182 were located 71 bp, 2358 bp, 37 bp, and 313 bp upstream of the 5' untranslated region, respectively. None of these P-element insertion lines were sensitive to caffeine in initial tests and they all complemented $huc95^{E}$ mutants for caffeine sensitivity. To confirm that the P-element insertions associated with these genes had affected gene function, RT-PCR of homozygous viable lines of P-element insertions associated with kal-1, CG6164, and CG6182^{Ey08283} was performed and revealed expected amplification products, suggesting that these genes were still being

transcribed in the homozygous lines. Since insertion of a P-element within the 5' untranslated region often inhibits or reduces gene function by interfering with transcription, the genes kal-1, CG6164, and CG6182^{Ey08283} cannot conclusively be eliminated as candidates. sec10 however, was not a homozygous viable P-insertion mutant and crosses to other deficiencies encompassing the 95D-E region also failed to produce viable hemizygotes, suggesting that the P-element insertion that created this mutant was a lethal insertion in that chromosome. Since the P-element only mapped 37 bp upstream of sec10 and had resulted in a lethal insertion, it is highly likely that this Pelement disrupts the function of this gene by inserting into the 5'-untranslated region. Furthermore, knockdown of Drosophila sec10 expression, which was previously done using a transgenic RNAi approach, resulted in post-embryonic lethality with larvae dying as second larval instars, suggesting that sec10 is an essential gene (Andrews et al., 2002). Since this P-insertion maps only within 37 bp of the upstream region (and thus is predicted to disrupt transcription and expression of sec10) and is associated with homozygous lethality, consistent with knockdown studies reducing Sec10 expression, this *P*-element insertion line can be considered a true *sec10* mutant and conclusively ruled out as a candidate of $huc95^{E}$ through complementation. Complementation tests were also done with several other *P*-element insertion lines $P\{EP\}CG6178^{EP3637}$, $P\{EP\}CG6178^{EP3251}, P\{EPgy2\}CG6178^{EY07693}$ and $P\{Sup \text{ or } P\}CG6178^{KG05318}, \text{ all of }$ which are associated with the gene CG6178, inserting 206 bp, 65 bp, 212 bp, and 214 bp upstream of this gene, respectively. All of these lines also complemented $huc95^{E}$ suggesting that this gene was not a candidate, or that the *P*-element insertions associated with this gene do not completely disrupt the function of this gene. To examine this

possibility, I performed RT-PCR on RNA isolated from these lines and was able to amplify CG6178 mRNA, suggesting that these *P*-elements do not knock out the function of this gene.

Section 2-3d: Sequencing of candidate genes

In addition to the strategies described earlier for determining which gene candidates could be associated with $huc95^{E}$, candidates whose known or proposed functions could plausibly explain the caffeine sensitive phenotype were sequenced and analyzed for sequence alterations. One attractive candidate was the Mpk2 gene (also known as p38a), which encodes a <u>M</u>itogen <u>A</u>ctivated <u>P</u>rotein <u>K</u>inase (MAPK) with homology to the mammalian MAPK p38. These kinases are known to have roles in response to various forms of cellular stress such as osmotic stress, inflammatory cytokines, presence of lipopolysaccharide, and UV irradiation (Seger and Krebs, 1995; Ono and Han, 2000; Johnson and Lapadat, 2002). The MAPK signal transduction pathway is conserved from yeast to mammals and consists of a complex cascade that involves activation of MAPK by MAPK kinases (MKK) that have been activated by MKK kinases (MAP3K) upon exposure to environmental stress, growth factors, cytokines or other mitogenic factors (Ono and Han, 2000). All p38 MAPKs are activated by dual phosphorylation from MKKs at a Thr-Gly-Tyr (TGY) motif and once activated, regulate the activity of transcription factors to control gene expression (Ono and Han, 2000). In mammalian systems, p38 MAPKs participate in innate immunity, inflammatory response, and stress-induced signalling events including a link with cell cycle checkpoint regulation. Previously, p38 was implicated in mediating a caffeineinsensitive (and thus an ATM/ATR independent) G2 delay following UV irradiation through phosphorylation of cdc25B and p53 (Bulavin et al., 2001; Goldstone et al., 2001; Bulavin et al., 1999). This suggests that p38 is involved in a separate checkpoint pathway that does not involve ATR or ATM (Bulavin et al., 2001; Bulavin et al., 2002). Because the premise of the original screen that identified $huc95^{E}$ was to discover potential checkpoint components, p38a seemed like an ideal candidate, as one could conceive a scenario whereby the MAPK pathway (which normally responds to environmental stress or pathogenic invasion) might also serve a back-up role in cell checkpoint control and cell cycle arrest. By this reasoning, $huc95^{E}$ flies might be sensitive to caffeine because they had lost this function, due to simultaneous loss of MAPK-mediated and ATR/ATM-mediated checkpoint arrests as a result of caffeine's inhibition of ATR/ATM (see Figure 2-8). To address this possibility, the coding region of Mpk2 was tested for any single base pair mismatches by annealing single strands of Mpk2 coding sequences amplified from $huc95^{E}$ to wild type genomic DNA and determining the annealing time (with the expectation that if there was a mutation, the annealing time would be altered) (Silva, 2002). By this test, no alterations appeared to exist within this region. To further test the possibility that alterations existed in the noncoding or regulatory regions of Mpk2, 5645 base pairs encompassing this entire gene including exons and the 5' and 3' untranslated regions (making up 2795 base pairs) were sequenced. No sequence alterations exist within this gene and its outlying regions, thus eliminating this gene as a candidate. Later, complementation tests were performed with a Mpk2 homozygous null mutant that was created by mobilization of a nearby P-element $P{EP}{CG6178^{EP3637}}$, 820 bp upstream of the Mpk2 locus (creating a new line containing a second P-element insertion 263 base pairs downstream of the Mpk2 locus), followed by

the simultaneous excision of both elements to create a precise 1.9 kb deletion that completely removed Mpk2 (Craig et al., 2004). RT-PCR analysis, as well as PCR of genomic DNA isolated from this stock, had confirmed the absence of Mpk2 and verified this mutant line. This homozygous null mutant failed to display any sensitivity to caffeine and was able to complement $huc95^{E}$ for caffeine sensitivity, confirming that Mpk2 was not responsible for the caffeine sensitivity of these mutants. Recently another putative gene, known as p38c with sequence homology to Mpk2, was also found to exist in the region immediately adjacent to the left of Mpk2. This gene was thought to arise as a result of a duplication event of Mpk2 (due to its sequence homology and proximity to *Mpk2*). However, *p38c* contains changes within its sequence at conserved motifs, which would interfere with its function as a MAPK, suggesting that p38c is actually a pseudogene. In particular, the invariant dual phosphorylation motif TGY was changed to a TDH by a replacement of the conserved tyrosine residue by a histidine in p38c (Craig, 2003). Also, changes within its catalytic loop, as well as the fact that no expressed sequence tags were found, suggested that p38c is likely a pseudogene. In case this putative MAPK did have some function that could be related to Mpk2 in other systems, the coding region of this gene was also sequenced in the $huc95^{E}$ hemizygotes and compared to wildtype sequences. This gene was also found to contain no sequence alterations within the coding regions, suggesting that this gene was not $huc95^{E}$. Because sequencing of the coding regions did not exclude the possibility that mutations might exist within the regulatory region, qualitative RT-PCR was performed to determine if the p38c gene was expressed. Results from this test showed that p38c is expressed, eliminating this putative gene as a candidate (see Figure 2-9). Although no expressed sequence tags have been reported thus far for this gene, this evidence for expression might be the result of the amplification process, which would serve to enhance undetectable amounts of RNA.

Another gene that was eliminated by genomic sequencing was *mRpS24*, localized at 95E6. This gene, which is predicted to encode a mitochondrial ribosomal protein, was easy to sequence because the reported size of this gene was only 718 bp with no introns. Furthermore, being a ribosomal protein localized in the mitochondrion (which is known to be linked to pathways involving programmed cell death), a scenario could be imagined whereby a mutation in this gene causes caffeine sensitivity that results in apoptosis. Sequencing results revealed no sequence alterations within the coding region of this gene and qualitative RT-PCR also showed that this gene was expressed in both wildtype and hemizygote flies, suggesting that this gene is not a candidate for the *huc*95^E mutation (Figure 2-9).

CG6178 is another candidate for sequencing that has a putative role in acyl-CoA biosynthesis (Oba *et al.*, 2004). CG6178 shares a high similarity to firefly luciferase ($e=8^{-113}$), a bifunctional enzyme that has monooxygenase (mediating bioluminescence) and a fatty acyl-CoA synthetase activity. *In vitro* catalytic tests have revealed that CG6178 does not exhibit luciferase activity, but is capable of synthesizing fatty acyl-CoA in the presence of ATP, Mg²⁺, and CoA (Oba *et al.*, 2004). Acyl-CoA and its derivatives can mediate cellular signalling by release Ca²⁺ from RyR channels and possibly stimulating Ca²⁺ reuptake at IP₃R channels (Faergeman and Knudsen, 1997; Fitzsimmons *et al.*, 2000). Because caffeine also evokes a Ca²⁺ release from RyR channels and could theoretically affect Ca²⁺ signalling from IP₃R channels (by affecting PKC signalling

through adenosine GPCRs), this gene was also a plausible candidate (Rousseau *et al.*, 1988; McPherson *et al.*, 1991). Upon sequencing 4743 bp for this gene, no sequence alterations were found, eliminating this gene as a candidate.

Other candidates that were sequenced and had displayed no sequence alterations were the genes CG31133 and CG6182, respectively (Table 2-2). Both of these genes have unknown functions and are not obvious candidates for $huc95^{E}$ other than residing within the deleted region. Nonetheless, opportunity arose for these genes to be sequenced and as a result, were successfully eliminated as candidates for $huc95^{E}$.

A gene encoding a novel class I myosin called *myo*95E was also a possible candidate even though it was not an obvious candidate in terms of being implicated in cell cycle checkpoint regulation. However, a mutation in a myosin gene could exhibit caffeine sensitivity, as myosins are often regulated by calcium containing molecules such as calmodulin. Myosins are molecular motors that transduce energy from ATP hydrolysis into mechanical movement along actin filaments. They are known to be involved in several different processes including cell locomotion, muscle contraction, cytokinesis, endo/exocytosis, vesicle and organelle transport, signal transduction, and establishment of polarity (Berg *et al.*, 2001; Krendel and Mooseker, 2005). There are currently 17 known classes of myosins, based upon whether they function as dimers or monomers, as well as particular domains and motifs that are related to specialized functions (Tzolovsky *et al.*, 2002). Structurally, all myosins are characterized by the presence of an N-terminal heavy chain that harbours a conserved 80 kDa catalytic domain containing an ATP-hydrolyzing domain and actin-binding sites, required for movement of the myosin (Berg *et al.*, 2001). The catalytic domain consists of three
conserved loops (the P-loop, switch-1 region, and switch-2 region, respectively) with consensus sequences that surround the active site (Yamashita et al., 2000). The conservation of these functional domains has been instrumental in the identification of other myosins, of which 13 have been identified in Drosophila (Yamashita et al., 2000; Berg et al., 2001; Tzolovsky et al., 2002). The P-loop serves as an ATP binding site, while the switch-1 region and switch-2 region interact together to hydrolyze ATP and transmit this energy beyond the active site (Smith and Rayment, 1996; Onishi et al., 1998; Furch et al., 1999). Following this N-terminal motor domain is an α -helical light chain, "neck" region that contains anywhere from 0 to 6 tandem isoleucine-glutamine (IQ) motifs, which serves as a binding site for regulatory proteins such as calmodulin or other members of the EF-hand containing family (Tzolovsky et al., 2002). This family of regulatory proteins is characterized by the presence of helix-loop-helix motifs that contain highly conserved residues in the loop region for the binding of Ca^{2+} (Kawasaki *et* al., 1998). It is thought that interaction of these regulatory proteins with Ca^{2+} affects the affinity of these proteins for IQ motifs in the regulatory neck region of the myosin. In general, unconventional myosins (which are all myosin classes that are not of the typical class II muscle myosins) bind calmodulin with higher affinity when it is not associated with Ca^{2+} (Tzolovsky *et al.*, 2002). This Ca^{2+} -sensitive binding of regulatory proteins to IQ motifs contributes to the regulation of these molecular motors, with the neck region postulated to act as a lever arm to amplify small movements in the motor domain to create the power stroke (Howard, 1997). In addition, the neck region of many myosins is the site of alternative splicing to produce myosin variants with different numbers of IQ motifs, adding further complexity to the regulation of these proteins. Following the neck region is the highly divergent C-terminal tail region, which has been known to contain various different motifs such as coiled-coil α -helical domains (eg. class II myosins) for dimerization of myosin monomers, Src homology 3 (SH3) domains (found in many intracellular or membrane-associated proteins that mediate diverse processes including the formation of multiprotein complexes, altering subcellular localization, or mediating the local accumulation of proteins), GAP domains, band **<u>F</u>** e_zrin-<u>r</u>adixin-<u>m</u>oesin homology (FERM) domains (which are involved in localizing proteins to the plasma membrane), Pleckstrin homology (PH) domains (which target proteins to the plasma and internal membranes through association with phosphoinositides), and Myosin tail homology (MyTH) domains, to name a few (Oliver et al., 1999). These diverse tail or cargo domains are thought to either target the myosin to different cellular localizations, to anchor the myosin thereby allowing it to move actin filaments within the cell, or to interact with particular cargo molecules and facilitate their transport to other locations by movement along actin filaments. Some tails also have kinase domains, PH domains or GTPase activating domains, suggesting that these myosins may also play a role in signal transduction.

By sequence homology and structural organization, *myo*95E encodes a class I myosin (Tzolovsky *et al.*, 2002). These were the first unconventional myosins (ie. non-muscle myosins) discovered and have been implicated in several processes including establishment and maintenance of cortical tension, motility, endocytosis and exocytosis (Osherov and May, 2000; Wu *et al.*, 2000). They lack a coiled-coil protein interaction domain within the tail region, suggesting that these myosins act as monomeric filaments. The tail also has a domain consisting of many basic amino acid residues referred to as the

Tail **h**omology **1** (TH1) region. This TH1 domain has been shown to bind phospholipid vesicles *in vitro* and is thought to play a role in tethering some class I myosins to membranes *in vivo* (Osherov and May, 2000). In addition to *myo*95E, *Drosophila* also contains 2 other class I myosin genes named *myo1A* and *myo1B*. *myo1A* and *myo1B* were discovered by a PCR-based hybridization approach, using degenerate primers to two highly conserved head domain regions (Morgan *et al.*, 1994). *myo1A* was localized to the cytological region of 31D-F and contains two IQ motifs, while *myo1B* was localized cytologically to 61F and contains three IQ motifs. Both of these class I myosins are associated with the brush border of the alimentary canal in both larval and adult stages, with Myo1A predominantly localizing to the sub-apical terminal web domain and Myo1B localizing to the apical microvillar domain (Morgan *et al.*, 1995). Like other brush border myosins in vertebrates, both Myo1A and Myo1B are thought to be involved in vesicle transport and regulation of membrane dynamics, possibly acting as structural elements within the brush border.

*myo*95E is a recently discovered class I myosin that was initially missed in the preliminary annotation of the *Drosophila* genome. Originally, the predicted sequence for this gene was unusually short for a myosin, encoding only 59 amino acids, which was later revealed to be the result of the gene being predicted incorrectly (Tzolovsky *et al.*, 2002). As a result, the sequence for this gene was manually assembled using expressed sequence tags (ESTs), the reported *Drosophila* genomic DNA sequence, and homology to other myosins. Two different splicing prediction programs theoretically predicted the splicing of this gene and the artificially assembled sequence was then used to design primers for RT-PCR amplification. The resulting cDNA was sequenced and it was

determined that the myo95E gene consists of 16 exons and produces at least 3 different types of transcripts that are expressed during oogenesis, larval, and adult stages. It was also determined by homology searches that the myosins produced from this gene share 33% identity (53% similarity) to vertebrate brush border class I myosins – even more so than to the two other class I myosins in Drosophila (Tzolovsky et al., 2002). myo95E also contains a 281 amino acid insertion within the N-terminal region, resulting from an unusually long exon 3, that is similar to an AAA (ATPase Associated with various cellular Activities) domain (Berg et al., 2001; Tzolovsky et al., 2002). Such domains exist in a wide variety of proteins in different organisms and are characterized by a conserved region of about 220 amino acids containing an ATPase domain that utilizes energy from ATP hydrolysis to carry out such processes as proteolysis, protein folding, membrane trafficking, cytoskeletal organization, organelle biogenesis, DNA replication, and intracellular motility (Hanson and Whiteheart, 2005). AAA motifs have also been identified in the molecular motor dynein, a microtubule-based motor required for chromosome movement, organelle transport, and ciliary/flagellar beating (Vale, 2000). AAA domains have not been previously found in myosins, which usually contain small motor domains with only one ATP binding site that are thought to modulate their activity by a different mechanism than dynein (Hanson and Whiteheart, 2005). It is believed that this domain modulates the hydrolysis of ATP as it is inserted into loop 1 of the motor domain, which harbours the other region for ATP hydrolysis. Myo95E has also been found to contain 2 IQ motifs with two different variants due to alternative splicing (Tzolovsky et al., 2002).

When I sequenced myo95E from genomic DNA isolated from $huc95^{E}$ hemizygotes, I found that the mutant allele contained a single nucleotide alteration that changed a guanine residue to an adenine residue. This mutation was confirmed in both the sense and anti-sense sequences, using two different primer sets on two different sequencing templates and is consistent with those commonly caused by EMS mutagenesis as methylation of G residues to O^6 -ethyl-G mispairs with T during replication, resulting in G/C to A/T transitions (Bentley et al., 2000). Analysis of this alteration revealed that this mutation existed at the junction of intron 8 and exon 9. mutating a normal AG splicing acceptor site to AA (see Figure 2-10A). This mutation identified in *my095E* is predicted to prevent splicing at this acceptor site, resulting in a severely truncated protein due to the presence of a TAA stop codon within the original reading frame 54 bp downstream of exon 8 in intron 8. Such a defect would result in loss of approximately half of the protein, including the terminal part of the motor domain, the entire neck domain (crucial for regulation) and the tail domain (crucial for function through interaction with specific cargo for transport or cellular components for anchoring). Consequently, the prediction is that this would result in a non-functional myosin. Furthermore, alternative splicing linking exon 8 to exon 10 (essentially bypassing the mutated splicing acceptor site at the junction of intron 8 and exon 9) would still result in a truncated protein that would be lacking the neck and tail regions due to a codon being formed in frame 233 stop bp in exon 10 with this processing (Figure 2-11A). Similarly, fusion of exon 8 to exon 11 would result in a myosin with no regulatory IQ motifs and splicing products eliminating exon 8 would also eliminate the switch-2 motif that is needed for ATP hydrolysis (Figure 211A). Only class XIV myosins do not contain any IQ motifs and are found only in Apicomplexan parasites such as *Toxoplasma gondii* and *Plasmodium falciparum* (Heintzelman & Schwartzman, 1997; Hettmann *et al*, 2000). Class XIV myosins are required to mediate a unique substrate-dependent gliding motility involving circular and forward twisting movements. Although it is not certain that a Myo95E isoform without IQ motifs would be nonfunctional, it is likely, as the only class of myosins lacking these motifs are involved in a specific type of movement for a single cellular organism. Furthermore, it could be possible that some other form of alternative splicing exists which does not use these typical splicing acceptor sites to result in a functional protein but based upon known acceptor sites, myosin functions, and sequence, myosins resulting from this mutation are predicted to be nonfunctional. In fact, all predicted full-length transcripts of myo95E (with the exception of one transcript which results in a severely truncated protein due to a stop codon in exon 5 and is thought to be inactive (Tzolovsky *et al.*, 2002)) utilize this particular splicing acceptor site, suggesting that if alternative splicing occurs, it does not involve this particular site (Figure2-11B).

To confirm that this splicing defect was actually present and determine if the myo95E gene was expressed, RT-PCR was performed on total RNA isolated from adult $huc95^{E}$ hemizygote flies and compared with third chromosome isogenized wild type flies using primers that flanked the intron (see Figure 2-10B). From this analysis, it was clear that hemizygotes were only capable of amplifying a 376 bp fragment (representing an unspliced transcript), whereas wild type controls displayed both the 376 bp fragment (in a stoichiometrically lower amount, possibly relating to the efficiency of the splicing reaction) and a 279 bp fragment (which resulted from the removal of a 97 bp intron by

splicing). These results implicated myo95E as a possible candidate for $huc95^{E}$, a hypothesis I pursued further by the following approach.

Section 2-3e: Transgenic rescue experiments

To test the possibility that mvo95E is the gene associated with the $huc95^{E}$ caffeine sensitivity phenotype, a genomic rescue plasmid containing the entire *myo*95E gene along with 2672 bp of the upstream and 1579 bp of the downstream regions was constructed utilizing the strategy outlined in Figure 2-12 and used to obtain transformants. Two putative transformants were obtained and the P transposon insertions associated with the two lines were subsequently mapped to the third chromosome. Since the $huc95^{E}$ mutation also existed on the third chromosome, crosses were set up with Df(3R)crb87-4as shown in Figure 2-13A, to produce recombinants that would contain both the transgenic rescue construct and the deficiency. Putative recombinants were selected on the basis of inheritance of w^+ gene (associated with the construct) and e (associated with the deficiency) over TM3 Sb, e and tested for their inability to complement Df(3R)crb87-4, thereby selecting against recombination events that linked the construct and the erecessive mutation without the deficiency. Recombinants were then crossed to huc95^E mutants and tested in the presence and absence of caffeine (Figure 2-13B). Unfortunately, the construct did not rescue the caffeine sensitivity of the $huc95^{E}$ mutants. There are two possibilities for this result: either myo95E is not the $huc95^E$ gene, or the rescue construct was either not functional or not expressed properly to mediate caffeine tolerance. To determine if the rescue construct was expressed, RT-PCR experiments were performed on RNA isolated from recombinants containing the rescue construct isolated in the absence of caffeine. The results of this experiment indicated that these flies exhibited a *myo*95E splicing pattern similar to that in wild type *myo*95E controls, suggesting that the construct was expressing *myo*95E mRNA normally (Figure 2-14). However, although the results suggest that the rescue construct is being properly spliced, it is unknown whether or not the RNA is being translated or expressed sufficiently in the cells necessary to mediate caffeine tolerance (if in fact *myo*95E corresponds to *huc*95^E).

Section 2-4: Discussion

At present, the identity of the gene responsible for caffeine sensitivity in $huc95^{E}$ mutants and the underlying mechanism involved, remains elusive. Out of an initial list of 23 genes. I have managed to narrow the number of remaining possible candidates to 9, with one (myo95E) showing particular promise as a candidate gene. I have determined though sequencing of genomic DNA isolated from hemizygous huc95^E mutants that myo95E has a single transition mutation of a G to an A within a AG splicing acceptor site at the junction of intron 8 and exon 9. This mutation is predicted to result in a truncated protein where the terminal portion of the head domain (important for the motor action and generation of power for myosin movement), the neck domain (required for regulation of the myosin), as well as the cargo domain (for interacting with the structures or components to be moved) as a consequence of a premature stop codon located 54 bp down from the splice site. Alternative splicing products using splicing acceptor sites of other known exons are also predicted to result in nonfunctional proteins. RT-PCR utilizing primers flanking the splice site (see Figure 2-10B) confirmed this splicing defect, demonstrating both that this gene was expressed and that hemizygotes bearing this mutation had lacked the proper splicing of this intron. Unfortunately, rescue experiments to confirm that this gene is associated with $huc95^{E}$ caffeine sensitivity have failed; even though RT-PCR experiments confirm expression of the construct and splicing within $huc95^{E}$ hemizygotes expressing the transgene. There are several possible reasons as to why the rescue construct might have failed to rescue the caffeine sensitivity of the $huc95^{E}$ mutants. One possibility is that the transgene could have inserted within a region of the chromosome that affected levels of its expression or function. Another possibility is that other regulatory elements required for sufficient *myo*95E expression were missing within the rescue construct. As such, position effects, or failure to include upstream or downstream elements that are crucial in the regulation of the expression of this gene could explain the failure of this construct to rescue the $huc95^{E}$ mutant phenotype.

Class I myosins have been implicated in vesicle transport and fusion to the plasma membrane, motility, and the establishment and maintenance of cortical tension (Osherov and May, 2000; Wu *et al.*, 2000). Myo95E in particular, shares higher similarity and identity (53% and 33%, respectively) with vertebrate brush border class I myosins compared to the two other class I myosins identified in *Drosophila* (Tzolovsky *et al.*, 2002). These similarities suggest a function for myo95E in exo or endocytic events and vesicle transport, as class I myosins within the brush border have been implicated in processes involving absorption of nutrients (Skowron *et al.*, 1998). Furthermore, recent findings in 3T3-L1 adipocytes have implicated a class I myosin, myo1C, in the fusion of vesicles containing the GLUT4 glucose transporter receptor, to the plasma membrane (Bose *et al.*, 2002). PI3K signalling was found to regulate this process as inhibition of PI3K prevented fusion of these vesicles (Bose *et al.*, 2002; Bose *et al.*, 2004). Interestingly, overexpression of myo1C was shown to mediate fusion of vesicles containing the GLUT4 receptor to the plasma membrane, even when PI3K was inhibited

(Bose et al., 2002; Bose et al., 2004). One of the many effects of caffeine is that it interferes with PI3K activity by inhibiting the p110 catalytic subunit of PI3K. Since PI3K is an important signalling molecule involved in processes such as apoptosis, cellular proliferation, and cell growth, perhaps the reason for caffeine sensitivity within the huc95^{DE} mutants is that caffeine-induced inhibition of PI3K interferes with these processes, which require $huc95^{E}$ function. If $huc95^{E}$ is myo95E, then a possible reason for the conditional lethality of $huc95^{E}$ mutants in the presence of caffeine could be that myo95E expression is required to overcome the caffeine-induced insult (analogous to the overexpression of myo1C overriding PI3K inhibition). Although myo95E is classified as a class I myosin because it contains a basic tail, it differs from all other class 1 myosins discovered to date in that it also contains a partial AAA domain within the head region of the protein (Tzolovsky et al., 2002). The significance of this domain (ie. whether it is a functional domain and what it is used for) remains unclear, however. Proteins containing such domains have been implicated in a wide variety of processes such as protein degradation, DNA replication, membrane fusion, and movement of microtubule motors. Perhaps this domain is crucial for the function of myo95E and (if myo95E is the gene associated with the *huc*95^E mutation) its ability to impart caffeine tolerance. In addition to this, several different isoforms based upon alternative splicing have been found for myo95E, which might suggest that this gene might be involved in producing several different functions of this class I myosin. Consequently, myo95E remains an attractive candidate for $huc95^{E}$.

Besides *myo*95E, the genes *CG13605*, *CG13607*, *CG6129*, *CG6164*, *CG5463*, *kal-1*, *CG5510* and *CG13606* are also possible candidates that have yet to be eliminated.

CG5510 might have putative roles in secretion and intracellular transport and could result in a caffeine sensitivity phenotype whereby the transport of a particular component or processes involved may be required to address the caffeine insult (similar to the conjecture of myo95E being a candidate). CG6129 is an attractive candidate because it has homology to Homo sapiens Centrosomal NIMA-related Kinase 2 (Nek2) Associated Protein 1 (C-Nap1) ($e=2.0^{e-27}$). The NIMA (<u>N</u>ever In <u>M</u>itosis in <u>Aspergillus nidulans</u>) protein kinases were originally discovered in the filamentous fungus Apergillus nidulans where they are required for regulating the onset of mitosis in collaboration with the cyclin dependent protein kinase Cdk1 (Oakley and Morris, 1983). Overexpression of NIMA causes premature mitotic events (such as chromatin condensation and formation of the mitotic spindle) and expression of Apergillus nidulans NIMA in Schizosaccharomyces pombe, Xenopus, and humans also induces premature mitotic events, suggesting that NIMA related pathways also exist in these organisms (Osmani and Ye, 1996). Nek2 is the most closely related mammalian homologue to NIMA and has been found to localize to the centrosome, the major microtubule organizing centre responsible for facilitating intracellular transport and imparting shape and polarity during interphase, as well as ensuring bipolarity and spindle orientation through formation of the spindle poles in mitosis (Fry et al., 1998a). Nek2 is also required for centrosomal splitting at the G₂-M transition, due to its ability to phosphorylate C-Nap1 (a core centrosomal protein that is responsible for holding the centrioles together during interphase), resulting in its displacement (Rapley et al., 2005; Fry et al., 1998a; Fry et al., 1998b). Interestingly Nek11, another related NIMA kinase, has been implicated in a DNA replication/damage and genotoxic stress checkpoint that exists downstream of a caffeine-sensitive pathway (Noguchi *et al.*, 2002). Because of homology to C-Nap1, its proposed role in cell cycle regulation, and the possibility that related proteins could exist in a caffeine sensitive pathway, CG6129 is an interesting candidate for *huc*95^E.

Amongst the other candidates, 3 genes (CG13607, CG6164, and CG13606, respectively) exist with no known homologues or putative function (see Table 2-2) and, as such, might be interesting candidates for possible novel functions, which may be potentially related to caffeine or cell cycle checkpoint control. The fastest way to determine the identity of $huc95^{E}$ is probably to sequence these remaining candidates as this would serve to directly confirm or eliminate these genes as possibilities. Also, isolation of more $huc95^{E}$ alleles could help to confirm any putative candidates by searching for mutations within these alleles. As such, efforts are continuing to identify $huc95^{E}$ by sequencing and a screen is being planned for the isolation of more alleles of this mutant. As a result, the identity of $huc95^{E}$ will not remain elusive for long.

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Gene name	Map position	Putative Function	Similar gene sequences		
CG10694	95E1	DNA repair protein	S. cerevisiae RAD23 (e=1.2e-21)		
syntaxin 1A	95E1	exocytosis	* syx1A alleles complement huc ^{95D-E}		
CG18428	95E1	unknown			
CG13605	95E1	protein ubiquitination			
CG13607	95E1	unknown			
CG6129	95E1-E3	Ciliary rootlet component	H. sapiens NEK2-assoc. protein (e=2.0e-27)		
CG5463	95E3-E4	tRNA amidotransferase	H. sapiens Gln-tRNA (e=1.6e-102)		
Tsc l	95E4	cell cycle regulator	* Tsc1 alleles complement huc ^{95D-E}		
sec10	95E4-E5	exocytosis	H. sapiens secretory protein (e=5.7e-148)		
CG6164	95E5	unknown			
kal-1	95E5	cell adhesion	H. sapiens anosmin-1 (e=4.9e-31)		
Mpk2	95E5	protein kinase	M. musculans p38 MAPK 14 (e=5.2e-134)		
p38c	95E5	putative protein kinase			
CG6178	95E5-E6	enzyme	P. pennsylvanica monooxygenase		
<i>myo</i> 95E	95E6	myosin ATPase			
CG6182	95E6	unknown			
mRpS24	95E6	ribosomal protein	<i>H. sapiens mRpS24</i> (e=3.4e-24)		
Apc2	95E6	microtubule function	* Apc2 ^{DeltaS} complements huc ^{95D-E}		
CG5510	95E6	glycoprotein	H. sapiens GP36B (e=1.7e-75)		
CG13606	95E7-E8	unknown			
CHORD	95E8	zinc binding			
CG5515	95E8	unknown	H. sapiens CGI-24 (e=4.9e-32)		
CG5524	95E8-F1	DNA repair protein	<i>S. pombe rad18</i> (e=1.6e-71)		
CG6204	95F1	unknown	S. pombe putative ATP binding protein $(z=6.02,52)$		
CCR4	95F1-F2	transcription factor	S. cerevisiae CCR4		
CG17786	95F2	unknown			

Table 2-1: Predicted genes in region 95DE of the third chromosome.

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Gene name	Map position	Putative function	<i>huc</i> 95 ^E candidacy
CG13605	95E1	Protein ubiquitination	?
CG13607	95E1	Unknown	?
CG6129	95E1-E3	ciliary rootlet component	?
CG5463	95E3-E4	glutamyl tRNA amidotransferase	?
Tsc 1	95E4	cell cycle regulator	Complements huc95 ^E
Sec10	95E4-E5	component of exocyst	Complements huc95 ^E
CG6164	95E5	Unknown	?
kal-1	95E5	homologue of cell adhesion protein anosmin-1	?
CG31133	95E5	Unknown	No sequence alterations
p38c	95E5	putative mitogen activated protein kinase	No sequence alterations
mpk2	95E5	mitogen activated protein kinase	No sequence alterations
CG6178	95E5-E6	acyl-CoA biosynthesis	No sequence alterations
myo95E	95E6	class I myosin	Putative splicing defect
CG6182	95E6	Unknown	No sequence alterations
mRpS24	95E6	mitochondrial ribosomal protein	No sequence alterations
Apc2	95E6	β-catenin chaperone protein	Complements huc95 ^E
CG5510	95E6	intracellular protein transport	?
CG13606	95E7-E8	Unknown	?

Table 2-2: Putative transcript	s that map to	the <i>Df(3R)W6</i>	region
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Stock genotype	Stock No.	Experiment	Donor
Df(3R)crb87-4	2362	Deficiency mapping Recombination	Bloomington stock center
Df(3R)crbF89-4	4432	Deficiency mapping Recombination	Bloomington stock center
Df(3R)mbc-R1	2585	Deficiency mapping	Bloomington stock center
Tp(3;2)slo3		Deficiency mapping	Bruce Ganetzky
Df(3R)W6	7251	Deficiency mapping	Marianne Bienz
$syx1A^{\Delta^{229}}$	4379	Recombination	Bloomington stock center
crb ²	3448	Recombination	Bloomington stock center
Tsc 1 ^{Q600X}		Complementation	Tian Xu
$Tsc l^{29}$		Complementation	Duojia Pan
Tsc I ^{Q87X}		Complementation	Iswar Hariharan
$Tsc I^{Q243X}$		Complementation	Iswar Hariharan
$Tsc I^{Q453X}$		Complementation	Iswar Hariharan
$Apc2^{\Delta S}$	5917	Complementation	Bloomington stock center
Apc2 ^{d40}	6801	Complementation	Bloomington stock center
Mpk2 ¹	8822	Complementation	Bloomington stock center
$Tp(2,3) ap^{Xa}, ap^{Xa}/CyO,$ $H\{w^{-mC} = P\Delta 23\} HoP2.1; Sb^{1}$	6394	P-element deletion synthesis	Bloomington stock center
ecd' , st' , red' , e^4 , ca'	4210	P-element deletion	Bloomington stock center
P{EP}Syx1A ^{EP3215}		synthesis P-element deletion synthesis	Szeged Drosophila collection
<i>P{EP}CG6178^{EP3637}</i>		P-element deletion	Szeged Drosophila collection
P{EP}mask ^{EP601}		P-element deletion synthesis	Szeged Drosophila collection
<i>P{GT1}CG33100^{BG01713}</i>	12631	P-element deletion synthesis	Bloomington stock center
w;Sp/CyO;Δ2-3 Sb,e/TM6, Ubx		P-element deletion synthesis	Scott Hanna
y ¹ ,w ^{67c23} ; P{w ^{-mC} y ^{-mDint2} =EPgy2} CG6182 ^{EY08283}	16885	Complementation	Bloomington stock center
w^{1118} ; $P\{w^{-mC}=XP\}$ kal-1 ^{d01966}	19168	Complementation	Bloomington stock center
w ¹¹¹⁸ ; PBac{w ^{mC} =WH} CG6164 ⁽⁰⁰³⁷⁹	18323	Complementation	Bloomington stock center
<i>P{EP}CG6178^{EP3251}</i>	17104	Complementation	Bloomington stock center
P{EPgy2}CG6178 ^{EY07693}	17388	Complementation	Bloomington stock center
<i>P{Sup or P}CG6178^{KG05318}</i>	13992	Complementation	Bloomington stock center

Table 2-3: Relevant fly stocks used in the mapping of $huc95^{E}$



Select non-TM3 Sb (Ser) as evidence of recombination under presence of caffeine

Figure 2-1. Strategy to map $huc95^{DE}$ through recombination. Crosses of the $huc95^{DE}$ mutation over a balancer containing TM3, Sb are crossed to stocks containing the recessive lethal mutants syx1A and crb in order to select for progeny flies containing both the $huc95^{DE}$ mutation and recessive lethal marker (by selecting against inheritance of the balancer). These flies are then crossed to a deficiency which uncovers both the $huc95^{DE}$ and recessive lethal mutations, and the progeny reared under caffeine, such that the only non-balancer containing flies that survive must have undergone a recombination event between the $huc95^{DE}$ and recessive lethal mutations. Note that caffeine treatment is not required for crosses involving the two recessive lethal mutations as these mutants are not caffeine sensitive.



Figure 2-2. Resultant recombination map of $huc95^{DE}$ mapping strategy. The positions of syx1A, crb, and $huc95^{DE}$ were determined through the recombination frequencies derived from crosses between combinations of double mutants with a deficiency that had uncovered all of these mutations. The distance between $huc95^{DE}$ and syx1A was determined out of a total of 3937 flies, and 5007 flies for $huc95^{DE}$ to crb. The distance between syx1A and crb was determined out of a total of 9541 flies.



Figure 2-3. *P*-element strategy for mapping *huc*95^{DE}. Males containing a $\Delta 2$ -3 transposase source element on the second chromosome (associated with the balancer CyO) and recessive markers (*e* and *ca*) to screen for recombinants on the third chromosome are generated and crossed to lines containing *P*-elements on the third chromosome within the vicinity of the gene to be mapped. The use of a translocation of the second and third chromosome marked with the dominant wing mutation apterous^{xasta} ensures inheritance of both the $\Delta 2$ -3 element and associated third chromosome (marked by *Sb*) together. Males containing the *P*-element, $\Delta 2$ -3 element, and recessive markers are then crossed to mutant females and the progeny screened for inheritance of only one of the recessive markers. Stocks of these recombinants are then constructed and crossed to the mutant in the presence of caffeine might harbour a flanking deletion, which extends into this gene region that can be mapped through molecular methods.







Α





CG13605-cont CG13605-text	CG13607-cont CG13607-text	C G 61 29-con C G 61 29-test T iic]-cont Tiic]-cont	CG5463-con) CG5463-test	CG6178-con CG6178-text	Apc2-cont Apc2-text	C G 5510- cont C G 5510- text	CG13606-cont CG13606-text
95E1	95E1	95E1-3 95E4	95E3-4	95E5-6	95E6	95E6	95E7-8



A ^{3R} chromosome



Figure 2-7: Localization of the $huc95^{DE}$ mutation to the cytological region of 95E1-E7-E8. A) The $huc95^{DE}$ mutation was mapped cytologically by the inability to complement Df(3R)crb87-4 and Df(3R)crbF89-4, and the ability to complement the synthetic deficiency Df(3R)slo3 (containing a deletion extending from 94D4-10 to 96A18, and the proximal portion of this deleted region (94D4-10 to 95E7-F1) duplicated, inverted, and inserted into the second chromosome at position 57B3-5). Further refinement was obtained by its inability to complement Df(3R)W6, which was later determined to contain the genes listed in **B**.



Figure 2-8: Possible model for a p38 mitogen activated protein kinase (MAPK) as a candidate for *huc*95^E. Because p38 MAPKs are involved in stress response and have been implicated in an ATM/ATR independent checkpoint pathway, a possible scenario for caffeine sensitivity is that p38 might be involved in an ATR/ATM independent pathway and can instigate a checkpoint when proper checkpoint regulation by ATM/ATR is compromised by caffeine. Hence, a mutation in p38 could render cells sensitive to damage if caffeine is used to inhibit ATM and ATR.







Figure 2-10: Splicing defect in *myo*95E in *huc*95^E hemizygotes. A) huc95^E hemizygotes exhibit a mutation within the eighth intron (in blue) in *myo*95E whereby a guanine at the intron 8-exon 9 junction is mutated to an adenine (shown in red), abolishing the proposed splicing acceptor site. Such a mutation would result in a severely truncated protein missing the terminal portion of the head region and the entire neck and tail region due to a premature stop codon (shown in green) along the original reading frame. B) Qualitative RT-PCR analysis demonstrating the splicing defect. RT-PCR was performed on total RNA isolated from an isogenized third chromosome stock used for the initial screen (lane 1) and from huc95^E hemizygotes using primers that flanked the intron (lane 2).

Α

Exon 8 translation:

VQETAQLLNMEAQILINCLTRANSTNSAQEDVGCEMDARQAATNRNTLCR TLYSRLFTWLVNKINESLKSTQREKNLALLDFYGFEALDHNSFEQFAINYS AEKIHQ

Exon 10 translation: reading frame from exon 9

IRHYASVVNYSIHRFLEKNSDMLPKYISAAFYQSKLSLVQSLFPEGNPRRQVT KKPSTLSSNIRTQLQTLLAIVKHRRSHYVFCIKPNEGKQPHQFDMALVQHQV RYMSLMPLVHLCRTGHCYHLLHVKFFHRYKLLNSLTWPHFHGGSQVEGIALI IRNLPLPSAEFTIGTKNVFVRSPRTVYELEQFRRLRISELAVLIQTMFRMYHAR KRFQRMRHSQMIISSAWRTWR

Exon 10 translation: reading frame from exon 8

DSSLCKCSELLNTSVSRKELRHAAEVHKRCLLSEQTFFGAKPIPRGESPS TGYQKAQHVEFEYPHPIADAAGHR*





Figure 2-11: Analysis of alternative splicing in myo95E that bypasses splicing of exon 9. A) The translated protein sequence of exon 8 is listed in red with the switch-2 motif shown in blue. Splicing of the intron between exon 9 and 10 would result in translation of exon 10 shown in green with the IQ motif shown in blue. Bypassing exon 9 results in a truncated protein as a result of a stop codon being formed (represented by the asterisk) in frame from translation of exon 8. B) The putative transcripts of myo95E. The red box designates intron 8, which is utilized in all of the putative full length transcripts (note that translation of myo95E-RG encounters a stop codon in exon 5 and also lacks the switch-1 and switch-2 motifs, probably making this truncated protein non-functional as a myosin.



Figure 2-12: Creation of myo95E genomic rescue construct. A 8039 bp fragment containing myo95E and most of the downstream genomic region was excised from BACR29F06 using *Scal* and *XhoI* restriction sites and cloned into *pBluescript* cut with *EcoRV* and *XhoI*. 3672 bp of the upstream genomic region was excised from BACR29F06 using *XhoI* restriction sites and cloned into the *pBluescript* containing the insert. Constructs containing the 3672 bp fragment ligated with the correct orientation were identified by PCR, and a 11711 bp fragment containing myo95E and upstream and downstream regions was excised using *EcoRI* and *XhoI*, and cloned into the transformation vector *pUAST*.

A

$$\frac{w}{Y}; \frac{Df(3R)crb87-4, e}{TM3. Sb} X \frac{w}{w}; \frac{P\{myo95E^{Rescue}, w^{+}\}}{TM3. Sb}$$

$$\frac{w}{w}; \frac{P\{myo95E^{Rescue}, w^{+}\}}{Df(3R)crb87-4, e} X \frac{w}{Y}; \frac{+}{TM3. Sb}$$

$$\frac{w}{Y}; \frac{P\{myo95E^{Rescue}, w^{+}\}}{TM3. Sb} X \frac{w}{W}; \frac{Df(3R)crb87-4, e}{TM3. Sb}$$

$$X \frac{w}{w}; \frac{Df(3R)W6}{TM3. Sb} X \frac{w}{Y}; \frac{TM3. Sb}{Non-TM3, Sb};$$
B
B

$$\frac{w}{Y}; \frac{P\{myo95E^{Rescue}, w^{+}\}}{TM3. Sb} \frac{Df(3R)crb87-4, e}{W}; \frac{huc95^{E}}{TM3. Sb}$$

$$E$$

$$\frac{w}{Y}; \frac{P\{myo95E^{Rescue}, w^{+}\}}{TM3. Sb} \frac{Df(3R)crb87-4, e}{W}; \frac{huc95^{E}}{TM3. Sb}}$$

$$E$$

$$\frac{w}{Y}; \frac{P\{myo95E^{Rescue}, w^{+}\}}{TM3. Sb} \frac{Df(3R)crb87-4, e}{WY}; \frac{huc95^{E}}{TM3. Sb}}$$

$$E$$

$$\frac{w}{Y}; \frac{P\{myo95E^{Rescue}, w^{+}\}}{TM3. Sb} \frac{Df(3R)crb87-4, e}{WY}; \frac{huc95^{E}}{TM3. Sb}}$$

$$E$$

$$\frac{W}{Y}; \frac{P\{myo95E^{Rescue}, w^{+}\}}{Lcaffeine}$$

$$\frac{W}{WY}; \frac{P\{myo95E^{Rescue}, w^{+}\}}{Lcaffeine}$$

Figure 2-13: myo95E transgenic rescue test for $huc95^{E}$ hemizygote caffeine sensitivity. A) To make an appropriate rescue line for the test, a recombinant stock was made containing both a deficiency that uncovered $huc95^{E}$ and the myo95E rescue construct and tested for lethality with deficiencies that uncovered $huc95^{E}$ to ensure integrity of the deficiency. Recombinants that fail to complement these deficiencies are crossed to $huc95^{E}$ mutants in the rescue test shown in **B**) in the presence and absence of caffeine, and the viability of hemizygotes carrying the genomic rescue construct assessed.



Figure 2-14: Hemizygotes expressing the rescue construct exhibit splicing of intron 8 in *myo*95E. Total RNA was isolated from huc95^E/ P{*myo*95E}, Df(3R)crb87-4, *e* and subjected to RT-PCR using primers that span the splice site of intron 8. Results are similar to RT-PCR performed with wild type flies and demonstrate splicing of intron 8 as compared to RNA isolated from hemizygotes.

Chapter 3: The Characterization of the Drosophila huc95^E Mutants

Section 3-1: Introduction

In addition to attempts at mapping and identifying the gene responsible for the $huc95^{E}$ mutation, some work was also done to further characterize these mutants. The initial identification and classification of the *huc* mutants was based upon the sensitivity of adult eye development to 2 mM caffeine and 3 mM HU treatment in a screen utilizing the EGUF-GMR-*hid* system for creating homozygous ommatidial clones (hence the name *huc* for <u>hydroxyu</u>rea and <u>caffeine</u> sensitive mutants) (Silva, 2002). When using this system, *huc*95^E mutants have compound eyes with severely reduced ommatidia that are occasionally fused and irregular (Figure 1-12). These mutants were subsequently found to be primarily sensitive to 2 mM caffeine, as treatment with 3 mM HU alone produced little discernible contribution to the phenotype (Silva, 2002). Furthermore, *huc*95^E hemizygotes survived to adulthood with no visible developmental abnormalities when grown on normal media, but undergo pupal lethality when grown on media containing caffeine concentrations as low as 0.5 mM. This suggests that *huc*95^E is a conditional mutant, implying that the *huc*95^E gene is not required for viability under normal growth conditions, but is necessary for development in the presence of caffeine.

The adult external structures emerge out of specialized, compartmentalized epithelial sacs called imaginal discs, which invaginate from the body wall during embryogenesis and proliferate rapidly without differentiating inside the larval body cavity (Cohen, 1993). These discs become spatially patterned along the body axis throughout larval development and upon pupariation (following the complete differentiation of these discs), they evert, resulting in the emergence of developed adult structures. The severe eye phenotype seen in the huc95E mutants prompted examination of the eye-antennal imaginal discs from EGUF/+; FRT82B huc95^E/FRT82B GMR-hid3R third instar larvae grown in both normal media and media containing HU and caffeine (Silva, 2002). The eye-antennal discs were smaller and more fragile in larvae reared in the presence of HU and caffeine as compared to larvae grown on normal media. To determine if there were any abnormalities in the differentiation of ommatidia in these mutants, these discs were stained with antibodies to a neuronal RNA-binding protein called embryonic lethal, abnormal visual (ELAV) (Figure 1-12) (Silva, 2002). This protein is required for normal maintenance and development of all neurons within the central nervous system, and interacts directly with neuronal RNA, such that ELAV antibody staining serves as a marker for those cells that have undergone neuronal differentiation (Yao et al., 1993). ELAV staining revealed that the differentiation of ommatidial precursors still initiates within these discs, with no change in the size of the individual pre-clusters (Figure 1-12). This suggests that the reduction in size of these discs is due to fewer, normal sized cells being present and not that the disc is composed of smaller cells (Silva, 2002). The pattern of pre-clusters also appeared to be disrupted, with clusters being more tightly packed and less evenly spaced as compared to heterozygous controls. This disorganization however, could be due to the fragility of the eye discs under HU and caffeine and thus, might be an artefact of dissection (Figure 1-12, compare C and E to D and F). To determine if there were any defects in the synchronization or coordination of cell cycles within the developing disc, the pattern of bromodeoxyuridine (BrdU) incorporation was examined in eye-antennal imaginal discs from EGUF/+; FRT82B huc95^E/FRT82B GMR-hid3R third instar larvae and compared to
eye imaginal discs from EGUF/+; FRT82B huc95^E/TM3 Ser third instar larvae in the presence of caffeine (Figure 1-13). BrdU is a base analog similar in structure to the nucleoside thymidine and is used to identify cells in S phase. Because cell cycles are synchronized at G₁ within the morphogenetic furrow, with unspecified cells undergoing a coordinated single division following the furrow (referred to as the second mitotic wave), BrdU staining could possibly reveal defects in cell cycle coordination. Homozygous mutant eye-antennal discs had fewer cells labelled with BrdU as compared to heterozygous controls, and had a staining pattern that was disrupted and more diffuse, suggesting that fewer cells are undergoing replication, or that cells are dying prematurely, or are arrested prior to undergoing mitosis or if the cells that did not undergo S-phase were arrested and capable of undergoing mitosis later on.

To further characterize the $huc95^{E}$ mutants, I performed experiments to analyze the sensitivity of these mutants to development in the presence of caffeine. Since the sensitivity of $huc95^{E}$ hemizygotes was only tested with caffeine concentrations as low as 0.5 mM, the approximate limit of caffeine sensitivity on the viability of $huc95^{E}$ hemizygotes was determined to test the acute sensitivity of these mutants. Also, since the intent of the screen that had isolated $huc95^{E}$ was to discover possibly novel cell cycle checkpoint mutants that would have remained hidden due to redundancy in checkpoint pathways, the effects of ionizing radiation on viability of $huc95^{E}$ hemizygotes was also examined. Furthermore, because $huc95^{E}$ hemizygotes die as pupal lethals in the presence of caffeine, the gene associated with this phenotype might have a temporal role in development (as the mutation does not immediately kill larvae grown in the presence of caffeine, but allows persistence to the pupal stage). The appearance of fewer ommatidia within mutants exposed to caffeine also suggested that cells within the eye imaginal disc that would eventually develop into ommatidia were unable to efficiently proliferate, or had undergone apoptosis, or both. As a result, experiments were done to look at the effects of caffeine treatment on cell proliferation and apoptosis during the development of the imaginal discs.

Section 3-2: Materials and Methods

Section 3-2a: Caffeine titration

Reciprocal crosses between $huc95^{E}/TM3$ Sb, Ser and Df(3R)crb87-4/TM3 Sb, Ser were set up in vials and bottles containing 0.5 mM, 0.25 mM, 0.125 mM, 0.06 mM, and 0 mM caffeine respectively, and the proportions of hemizygotes determined from the progeny of these crosses. Caffeine-containing food was prepared by scoring the media in vials and bottles with a clean metal spatula and adding 250 µl or 2.5 ml respectively, of an appropriate concentration of caffeine (dissolved in distilled water) to yield the desired concentration.

Section 3-2b: Ionizing Irradiation Sensitivity Assay

Reciprocal crosses consisting of $huc95^{E}/TM3$, Ser-GFP and Df(3R)crb87-4/TM3 Ser-GFP, were set up and allowed to develop to third instar larvae (to ensure that the larvae were as robust and mature as they could be in order to survive the irradiation treatment). The larvae were exposed for 0, 45, 90, 180, or 360 seconds in a cobalt⁶⁰ gamma source, to mediate exposure to 0, 5, 10, 20 and 40 Grays (Gy) of ionizing radiation, respectively. The crosses were then left to develop at 22°C and the progeny sorted and tabulated following eclosion.

Section 3-2c: Imaginal Disc Antibody staining

Third instar wandering larvae of an appropriate genotype were dissected in 0.01 M phosphate-buffered saline (PBS) and fixed in 3.7% formaldehyde, 0.1% polyoxyethylene sorbitan monolaurate (tween 20), and 0.01 M PBS (PBST) for 20 minutes. The discs were then rinsed three times in PBST for 5 minutes and blocked in 5% bovine serum albumin (BSA) in PBST for 30 minutes. Primary antibodies of choice at the appropriate dilutions were then added to discs soaking in 5% BSA in 1% PBST and incubated with gentle agitation on a rotating nutator at 4°C overnight. Following three 10-minute rinses in PBST, the discs were then incubated in the appropriate secondary antibodies at a dilution of 1 in 1000, in 5% BSA with no light at room temperature for 1 hour. Samples were then rinsed again in PBST three times for 10 minutes and stained with hoechst dye for 5 minutes at a dilution of 1 in 500. Following a final rinse for 10 minutes in PBST, imaginal discs or brains were then dissected out from inverted larval cuticles in PBS and mounted on microscope slides with 80% glycerol in PBS. The dilution for anti-mouse Phospho-Histone 3 (Ser10) antibody (Upstate Biotechnology Cat No. 05-598) was 1/1000, while the dilution for anti-rabbit cleaved human caspase-3 antibody (Cell signaling Cat No. 9661) was 1/800.

Section 3-3: Results

Section 3-3a: The viability of huc95^E hemizygotes is compromised at concentrations as low as 0.25 mM of caffeine

Originally, the viability of $huc95^{E}$ hemizygotes was found to be sensitive to 0.5 mM caffeine, but the limits of this sensitivity were not determined (Silva, 2002). A caffeine titration experiment was performed to test sensitivity of hemizygotes to 0.25

mM, 0.125 mM, and 0.06mM caffeine (Figure 3-1) (Krause, 2003). At concentrations of 0.25 mM, the proportion of hemizygote survivors was similar to concentrations of 0.5 mM for both reciprocal crosses. At a caffeine concentration of 0.125 mM, the proportion of survivors drastically increases to approximately 9 times the proportion seen at a concentration of 0.25 mM in both reciprocal crosses, and is comparable to proportions seen at 0.06 mM caffeine. Proportions exhibit only a modest increase when no caffeine is added as compared to 0.06 mM caffeine in both reciprocal crosses, suggesting that the caffeine concentration sufficient to compromise the viability of $huc95^{E}$ hemizygotes is between 0.25 mM and 0.125 mM caffeine.

Section 3-3b: Ionizing radiation does not substantially affect the viability of $huc95^{E}$ hemizygotes

Because the original premise of the screen was to isolate potential cell cycle checkpoint mutants, it was logical to determine if the $huc95^{E}$ mutants were defective in cell cycle checkpoint regulation. Since it was previously known that caffeine affected the viability of $huc95^{E}$ hemizygotes, it was reasoned that if $huc95^{E}$ was defective for a cell cycle checkpoint function, $huc95^{E}$ hemizygotes might also exhibit increased sensitivity to DNA damaging agents. Ionizing radiation is an effective inducer of DNA checkpoint activation and has been commonly used to study the effects of cell cycle checkpoint response by mediating DNA damage such as double stranded breaks (Iliakis *et al*, 2003). As such, the sensitivity of hemizygotes to ionizing radiation was assessed to determine if the $huc95^{E}$ mutation rendered it checkpoint defective. To test this, crosses and reciprocal crosses of Df(3R)crb87-4/TM3 Ser-GFP and $huc95^{E}/TM3$ Ser-GFP were set up and third instar larvae resulting from the cross were exposed to varying amounts of gamma

irradiation at 0, 5, 10, 20, and 40 Gy, respectively. After irradiation, the larvae were allowed to develop, and the proportion of hemizygotes determined (Figure 3-2). No progeny developed to adulthood when exposed to 40 Grays delineating this dosage as completely lethal, even to wildtype. Exposure to 20 Grays drastically reduced the total number of progeny that developed to adults and corresponded to a developmental delay of 3 days, which was not observed with lower dosages. There was no appreciable reduction in the number of hemizygotes that developed following gamma irradiation compared to unirradiated controls at dosages of 5 or 10 Gy, suggesting that hemizygotes are not sensitive to ionizing radiation. At 20 Gy, the proportion of hemizygotes does significantly decrease (with proportion levels approximately half of those at 10 Gy) however, the total number of flies was also reduced, suggesting that at this dosage, the viability of all the flies are affected, not just hemizygotes. Furthermore, *huc*95^E hemizygotes did not exhibit additional delays in development relative to their heterozygous siblings, further supporting the conclusion that these flies were not sensitive to ionizing irradiation.

Section 3-2c: $huc95^E$ hemizygote third instar larvae display increased apoptosis in eye imaginal discs when reared in media containing 2 mM caffeine

To determine if the reduction in ommatidial cells and overall size of the eye observed in EGUF/+; FRT82B huc95^E/FRT82B GMR-hid3R mutants was due to caffeine-induced apoptosis, eye discs from EGUF/+; FRT 82B huc95^E/FRT82B GMR-hid3R larvae grown in the presence and absence of 2 mM caffeine, were stained with acridine orange to visualize cell death (Bradley, 2002). The results revealed more cell death towards the anterior of the morphogenetic furrow in caffeine-treated larvae as

compared to larvae grown in the absence of caffeine, suggesting that more cell death is occurring in larvae exposed to caffeine (Figure 3-3). Because the larvae contained GMRhid (a construct containing the apoptosis-triggering gene <u>h</u>ead <u>involution defective</u> fused to the *Glass Multiple Reporter* promoter) which is expressed mainly in post-mitotic cells after the second mitotic wave, significant background staining was apparent posterior of the morphogenetic furrow (where *GMR* is active). To examine apoptosis without *GMR*hid, huc95^E hemizygous larvae were grown in the presence and absence of caffeine, and dissected for eye-antennal imaginal disc analysis. Third instar huc95^E hemizygous larvae grown on caffeine-containing media had either no imaginal discs of any kind, or discs so small and fragile that they escaped detection. Furthermore, huc95^E hemizygotes died as pupal lethals with no apparent development of any adult structures within the pupal cases, consistent with my observation that these larvae had no imaginal discs. This suggests that the caffeine-induced lethality and sensitivity of huc95^E mutants is due to the loss of imaginal discs, which would eventually differentiate and evert to form developed adult structures.

When testing $huc95^{E}$ hemizygous larvae in media containing caffeine, the caffeine was added to the fly media such that the embryos laid would be developing under the constant presence of caffeine. To determine if prolonged development in the presence of caffeine was necessary to produce these effects, I set up multiple vial crosses to obtain hemizygotes and subsequently added 2 mM caffeine to each vial 1 to 6 days after egg laying (AEL). Imaginal discs from hemizygotes were obtained after addition at days 5 to 6 in the presence of caffeine, corresponding to addition of caffeine at the third larval instar. This suggests that caffeine might have a role in either the prevention of the

formation of imaginal discs, or its destruction. If caffeine did result in degeneration of the imaginal discs, this might explain the increased acridine orange staining that was attained with EGUF/+; FRT82B, huc95^{DE}/FRT82B, GMR-hid3R larvae grown in the presence and absence of 2 mM caffeine. Another observation is that there was always a developmental delay of about 3 days after addition of caffeine, which is not surprising since toxic components or stress result in developmental delays. To see if caffeine was causing the destruction of imaginal discs, 2 mM caffeine was added after day 4, and the larvae were allowed to develop to the wandering third instar larval stage. Dissection of these larvae revealed no imaginal discs, however upon dissection of the larval brains and staining with an antibody to activated caspase 3, intense staining could be seen within tissues that would have resulted in imaginal leg and eye-antennal discs, suggesting that there were discs, but that they were in the process of deteriorating, thus becoming fragile and undetectable. (see Figure 3-4). The deterioration of these discs is accompanied by apoptosis, which might be a reason for the lack of imaginal discs and pupal lethality of these mutants. To further examine this possibility, 2 mM caffeine was added to the media after 4 days of development, and the larvae allowed to develop for an additional 2 days, after which, the larvae were dissected and imaginal eye-antennal and wing discs were stained with an antibody against phospho-histone-3 (PH-3) a histone protein that is phosphorylated in cells undergoing mitosis) and activated caspase 3 (see Figure 3-5 and Figure 3-6). When grown on normal media, $huc95^{E}$ hemizygotes exhibited very little caspase-3 staining that was comparable to heterozygous controls reared on normal media (Figure 3-5 and Figure 3-6: compare panel A and I). In the presence of caffeine however, imaginal discs from $huc95^{E}$ hemizygous larvae exhibited markedly more caspase-3

staining compared to heterozygous controls (which appeared only slightly increased relative to controls raised on normal media). Interestingly, imaginal discs dissected from heterozygous larvae also displayed a slight increase in caspase-3 staining (Figure 3-5 and Figure 3-6: compare panel I and M), however this staining was no where near the intensity of staining exhibited by imaginal discs from *huc*95^E hemizygous larvae reared in caffeine-containing media. Staining with PH-3 did not seem to show any differences in both huc95^E hemizygotes or heterozygous controls in either the presence or absence of caffeine. All imaginal discs in all conditions had exhibited a lot of PH-3 staining (consistent with the proliferative nature of third instar imaginal discs) conditions. Although PH-3 staining might appears to be different (ie. less abundant and more punctate) in the caffeine treated heterozygous controls and hemizygotes reared on normal media, this is believed to be an artefact of the staining procedure. As a result, these observations are preliminary and future work to quantify the number of PH-3 staining cells within better-dissected discs.

Section 3-4: Discussion

From this analysis, I determined that the viability of the $huc95^{E}$ mutants raised in the absence of caffeine is not sensitive to ionizing radiation treatment implying that $huc95^{E}$ is not likely to have a novel cell cycle checkpoint function. $huc95^{E}$ cannot be conclusively ruled out as a novel cell cycle checkpoint mutant however, as any roles pertaining to cell cycle checkpoint regulation might be masked by redundant mechanisms that compensate for $huc95^{E}$. Alternatively, it also could be possible that $huc95^{E}$ encodes a checkpoint component that is not involved in responses to ionizing radiation or DNA repair. $huc95^{E}$ mutants are sensitive, however, to caffeine concentrations of down to 0.25 mM. In the presence of caffeine, hemizygotes die at the pupal stage and examination of the pupal lethals revealed no evidence of adult structures. Furthermore, examination of third instar hemizygous larvae reared in the presence of caffeine revealed no imaginal discs, suggesting that the pupal lethality observed in these mutants was due to the inability to complete metamorphosis. I determined that this inability to develop imaginal discs could be relieved if caffeine was added to the media at day 5 AEL. Upon staining third instar larval brains with caspase-3 from hemizygotes grown in media that had caffeine added at day 4, intense staining of tissues that would eventually result in imaginal discs was seen, suggesting that day 4-5 was the critical time period for exposure to caffeine. Furthermore, imaginal discs dissected from $huc95^{E}$ hemizygotes grown in the presence of caffeine exhibit more caspase-3 staining compared to heterozygous controls and hemizygous controls reared in the absence of caffeine, indicating that the imaginal discs are deteriorating in the hemizygotes as a result of caffeine-induced apoptosis. No detectable changes were observed upon the staining of imaginal discs with the mitotic marker PH-3 in both $huc95^{E}$ hemizygous larvae and heterozygous controls both reared in the presence and absence of caffeine, suggesting that the process of mitosis remains unaffected. This is just a preliminary observation however, and further work to quantify the number of cells undergoing mitosis in both hemizygous and heterozygous controls would serve to confirm this result.

Although it is clear that $huc95^{E}$ mutants exhibit increased apoptosis upon exposure to caffeine, it is still unknown as to why and how caffeine induces cell death in these mutants. As a result, an interesting question to address is whether this occurs through the release of calcium. Calcium is an important signalling molecule and is routinely kept at an intracellular concentration of approximately 100 nM in resting cells (Hanson et al., 2004; Saris and Carafoli, 2005; Waring, 2005). Such concentrations are accomplished by controlling the entry or efflux of Ca^{2+} into, and out of the cell, and the active sequestration and storage of intracellular Ca^{2+} within the endoplasmic reticulum (ER) and mitochondria through various channels and transporters. Excessive or inappropriate calcium release that depletes intracellular Ca²⁺ ER stores, can lead to apoptotic events by triggering mitochondrial-mediated pathways of caspase activation and apoptosis through the release of cytochrome C into the cytoplasm (Hanson et al., 2004). As a result, it would be interesting to stain imaginal discs from $huc95^{E}$ hemizygotes exposed to caffeine with antibodies to cytochrome C and see if cytochrome C release is involved. Furthermore, RyRs (receptors that reside in the ER and play a role in the regulation of cytosolic Ca^{2+} are sensitive to caffeine and evoke Ca^{2+} release upon exposure (Rousseau et al., 1988; McPherson et al., 1991). Treatment of Chinese hamster ovary cells with caffeine and ryanodine induces a release of calcium from the ER through its actions on RyR, resulting in apoptosis as ER stores of Ca^{2+} are depleted (Pan *et al.*, 2000). Thus, it would be interesting to see the reaction of $huc95^{E}$ mutants to ryanodine. Drosophila has only one RyR gene (Ryr44F) but mutants in Ryr44F exhibit impaired muscle function and die as first larval instars (Sullivan et al., 2000). However, it would be interesting to see whether any interaction between Ryr and $huc95^{E}$ occurs by making a double mutant expressing a homozygous rvr mutant gene (using the EGUF-GMR-hid system) and hemizygous $huc95^{E}$ in the absence of caffeine. Interestingly, genetic interaction studies in Drosophila involving Ryr44F and Cam (a gene located cytologically at 48F1 which encodes the Ca²⁺ sensor protein calmodulin) showed that the

one copy of a hypomorphic Ryr allele was capable of partially rescuing phenotypes mediated by the Cam^7 allele, which comprised of pupal lethality, pupal cases with deep indentations at larval segmental boundaries, and pharate adults with inverted heads inside the thorax (Wang *et al.*, 2003). Furthermore, the inhibition of phototransduction in *Drosophila* photoreceptors, mediated by treatment with ryanodine and caffeine, was rescued by the subsequent application of Ca²⁺-calmodulin (Arnon *et al.*, 1997). These results suggest that calmodulin plays a role in RyR regulation and thus, it would be interesting to determine if *Cam* and *huc*95^E interact and whether the *Cam* mutation could partially rescue the *huc*95^E mutation in the presence of caffeine. Alternatively, experiments involving Ca²⁺ channel blockers may also serve to elucidate the role of calcium channels in the apoptosis response of these mutants.

Another possibility is caffeine's effect on the phosphoinositide 3-Kinase (PI3K): an important signalling enzyme that has been implicated in a wide variety of processes ranging from cellular proliferation, apoptosis, differentiation, and metabolic regulation (Foukas *et al.*, 2002; Katso, *et al.*, 2001; Osaki *et al.*, 2004; Franke *et al.*, 2003). One of the most well studied aspects of PI3K signal transduction is its role in the insulinsignalling pathway (Shepherd, 2005). The hormone insulin rapidly stimulates class I_A PI3K signalling, which regulates such events as glycogen synthesis, inhibition of lipolysis, stimulation of protein biosynthetic pathways, and the translocation of the GLUT4 glucose transporter (a receptor required for the intake of extracellular glucose in to the cell) to the plasma membrane (Shepherd, 2005; Bose *et al.*, 2002). In addition to regulating glucose metabolism, insulin signalling is also the principle regulator of cell growth and development in *Drosophila* and is highly conserved from *Drosophila* to humans (Lizcano *et al.*, 2003; Bikopoulos *et al.*, 2004). *Drosophila* have one insulin receptor (DinR), seven insulin-like peptides, and several homologues of mammalian insulin receptor substrates such as PI3K, PTEN, PKB/Akt, and p70SK (Bikopoulos *et al.*, 2004). Expression of <u>Drosophila</u> insulin-like peptide-2 (DILP-2) results in an increase in body mass and growth in the developing eye when exclusively overexpressed in this tissue and inhibition of insulin signalling corresponds to a decrease in the overall size of the fly and a reduction in the development of imaginal discs (Edgar, 1999; Brogiolo *et al.*, 2001). These results reflect the ability of insulin signalling to inhibit apoptosis, thereby allowing continued growth of cells and tissues by preventing their destruction (Bertrand *et al.*, 1998; Lee-Kwon *et al.*, 1998; Yenush *et al.*, 1998).

Caffeine has been found to inhibit the lipid kinase activity of class I PI3Ks *in vitro* and block the ability of insulin to stimulate PKB (an important kinase that inhibits processes of apoptosis and regulates cell size) *in vivo* in CHO-IR cells and rat soleus muscle with IC₅₀ values similar to those required for inhibition of PI3K activity (Foukas *et al.*, 2002). This suggests that caffeine can inhibit insulin signalling by inhibiting PI3K and could explain the reduced size of imaginal discs and the increased occurrence of apoptosis in the *huc*95^E hemizygotes grown in caffeine since insulin signalling regulates cell size and apoptotic events. Furthermore, apoptosis has also been consistently linked with a decrease in glucose transport (Moley and Mueckler, 2000). Glucose transport and metabolism can trigger apoptosis because decreased glucose uptake results in ATP depletion, which can stimulate the mitochondrial cell death cascade or can result in oxidative stress, which can trigger stress-activated and other signal transduction pathways that signal apoptosis (Moley and Mueckler, 2000). Alternatively, intracellular

hypoglycemia and hypoxia due to decreased glucose transport can increase expression of **h**ypoxia-**i**nducible **f**actor-**1** α (HIF-1 α), which stabilizes p53 and activates p53-associated apoptotic response pathways (Moley and Mueckler, 2000). Because caffeine also blocks insulin-stimulated glucose transport in CHO-IR cells, caffeine could conceivably signal apoptosis by affecting glucose uptake and metabolism (Foukas *et al.*, 2002).

Upon stimulation by insulin, the class I myosin Myo1C is required to facilitate the fusion and allocation of vesicles containing GLUT4 receptors to the plasma membrane in 3T3-L1 adipocyte cell cultures, and inhibition of PI3K by the inhibitor LY294002 could block this fusion step, resulting in GLUT4-containing vesicles accumulating just beneath the plasma membrane (Bose et al., 2002; Bose et al., 2004). This block in fusion mediated by PI3K inhibition could be relieved however, by overexpression of Myo1C (Bose et al., 2004). The Drosophila class I myosin myo95E exhibits a polymorphism suggestive of a splicing defect that is predicted to result in a non-functional protein in the huc95^E mutants. Structurally, Myo1C differs from Myo95E in that it contains three IQ motifs and a cargo domain that contains several proline-rich domains that are thought to interact with SH3 domains. In BLAST alignments however, Myo95E is more related to Myo1C than to the other two class I myosins within Drosophila, suggesting that these differences may not be relevant (Tzolovsky et al., 2002). Accordingly, an explanation for caffeine sensitivity of the $huc95^{E}$ mutation (if myo95E is the gene associated with the $huc95^{E}$ mutation) is that the expression of myo95E is required to compensate for caffeine's inhibition of PI3K signalling (analogous to overexpression of Myo1C being required to overcome a block in fusion mediated by PI3K inhibition in cells). For instance, a possible model could be that caffeine's ability to inhibit PI3K signalling results in a decrease in downstream signalling mediated by PIP₃, including the activation of PKB (which renders cells extra sensitive to caffeine's ability to signal apoptosis), and inhibition of glucose transport (which triggers apoptosis to occur) (Figure 3-7). This results in degeneration of the imaginal discs and the consequent pupal-lethality (as no imaginal discs would be present to allow further metamorphosis). Expression of *myo*95E may be required to alleviate this sensitivity, possibly by allowing for vesicular trafficking to continue and inhibit apoptosis from occurring by maintaining glucose transport. This might also suggest why the rescue construct was not successful in rescuing the caffeine sensitive phenotype as expression of the construct may be reduced to that of wildtype.

To test this model, it will be necessary to determine if caffeine is capable of inhibiting PI3K signalling in *Drosophila*. PI3K signalling also regulates autophagy: a conserved process whereby amino acids and other nutrients from long-lived proteins, cellular organelles and other components are recycled within autophagic lysosomes (Rusten *et al.*, 2004). The process of autophagy is generally activated in response to environmental stress and nutrient deprivation and sustains the viability of cells under suboptimal conditions by recycling non-essential macromolecules (Klionsky and Emr, 2000). In holometabolous insects, autophagy is also triggered temporally in conjunction with development and metamorphosis, as larval tissues and organs are degraded and reorganized into adult forming structures. In this context, autophagy occurs by compartmentalization of cytoplasmic components by a cisternal isolation membrane derived from a preautophagosomal structure (PAS) to form a double membrane-enclosed autophagosome (see Figure 3-8) (Baehrecke, 2002; Noda *et al.*, 2002). Fusion of the autophagosome with a lysosome forms an autolysosome: the site of degradation where

cytoplasmic components are recycled (Fengsrud et al., 2000). The mechanisms underlying formation of the PAS have been elucidated from studies in yeast and involve two ubiquitin-like conjugation systems. Initially, a small ubiquitin-like protein Atg12, covalently attaches to another ubiquitin-like protein Atg5, to form an Atg12-Atg5 conjugate (Mizushima et al., 1998). This conjugate requires another ubiquitin-like protein Atg8, conjugated to the phosphatidylethanolamine (PE), for localization to the PAS (Suzuki et al., 2001). Both conjugates are required for the process of autophagy to occur and ultrastructure analysis has revealed that Atg5 and Atg8 (or its mammalian homologue Microtubule associated protein-1 light chain $\underline{3}$ (LC3)) localize to the isolation membranes (Mizushima et al., 2001). Upon closure however, Atg5 is lost while LC3 still remains, making LC3 a marker for newly formed autophagosomes. Activation of the insulin pathway, and the consequent stimulation of class I_A PI3K signalling, represses autophagy in mammalian cells and Caenorhabditis elegans, suggesting that PI3K has a negative role in the regulation of autophagy (Melendez et al., 2003; Seglen and Bohley, 1992). This is consistent with insulin's role as a regulator of cell growth and apoptosis as the larval period is characterized by an increase in mass with growth occurring predominantly in larval-specific tissues such as the salivary gland, epidermis, musculature, and fat body. At the end of the larval stage, these tissues and organs are marked for developmentally programmed autophagy to make way for the formation of adult structures. The fat body is primarily capable of starvation-induced autophagy and acts as a sensor of nutritional status to maintain adequate nutrient levels within the haemolymph and ensure survival during unfavourable conditions. Larval starvation results in the rapid inactivation of PI3K yielding reduced levels of PIP₃ in the plasma membrane of fat body cells. Mutations disrupting PI3K phenocopy many of the effects of starvation, including reduced larval growth, specific arrest of endoreplicative cell cycles, and aggregation of lipid vesicles within the fat body, suggesting a role for PI3K in repressing autophagy (Britton et al., 2002; Colombani et al., 2003). Alternatively, constitutive expression of PI3K within the fat body prevents these responses and results in reduced viability under starvation conditions, suggesting that autophagy of the fat body serves to provide adequate nutrient levels during times of stress (Britton et al., 2002). The process of starvation-induced autophagy was recently studied using a technique whereby stages of autophagy can be visualized utilizing transgenic lines harbouring a UAS-LC3-GFP or UAS-Atg5-GFP construct and a GAL4 driver specifically expressed within the haemolymph or fat body (Rusten et al., 2004). This technique, along with use of the vital dye lysotracker (which stains acidic vesicles), is capable of identifying three distinct compartments representing three populations within the autophagy pathway: autophagosomes (which contain Atg5 or Atg8 (LC3) and thus could be visualized through GFP staining), lysosomes (stained with lysotracker), and autolysosomes (which would involve vesicles that co-stained with both GFP and lysotracker). Utilizing this technique, premature autophagy was demonstrated on second and third instar feeding larvae (previously shown not to exhibit the effects of autophagy) that were starved for 4 hours on amino acid deficient sucrose medium (as evident from co-staining and the presence of larger, more acidic autolysosomes) as well as by treatment with the hormone ecdysone. Expression of dp110 (the catalytic subunit of PI3K) or dAkt (one of the substrates of PI3K signalling) also resulted in a strong reduction of GFP-LC3/lyostracker positive structures (in a similar manner to results involving ecdysone expression), and analysis of clones expressing of a dominant negative

ecdysone receptor had displayed higher levels of PI3K activity compared to neighbouring cells had correspondingly fewer and smaller acidic staining vesicles. Hence, this technique can provide a unique means by which one can determine if exposure to caffeine is capable of inhibiting the PI3K pathway *in vivo* in *Drosophila* larvae, by assaying the process of autophagy in the presence of caffeine.

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Figure 3-1: Caffeine sensitivity of $huc95^{E}$ hemizygotes. Reciprocal crosses were set up between $huc95^{E}/TM3$ Sb, Ser and Df(3R)crb87-4/TM3 Sb, Ser in fly media containing the caffeine concentrations listed. The proportion (listed above the bars) of $huc95^{E}$ hemizygotes that survived to adulthood was determined from the total progeny (listed in white in the bars)



Radiation exposure

Figure 3-2: Effects of ionizing radiation on huc95^E hemizygote viability. Reciprocal crosses were set up between $huc95^{E}$ / TM3 Ser-GFP and Df(3R)crb87-4/ TM3 Ser-GFP, allowed to develop to third instar larvae, and irradiated at the exposures specified using a Co⁶⁰ gamma source. The proportion (listed above the bars) of $huc95^{E}$ hemizygotes that survived to adulthood was determined from the total progeny (listed in white in the bars)



Figure 3-3: Eye discs dissected from EGUF/+; FRT82B huc95^{DE}/ FRT82B GMRhid3R CL3R larvae that were stained to visualize apoptosis by acridine orange staining. Disc from a larva grown on normal media (A) or from larvae grown on caffeine-containing media (B). Apoptosis is elevated in cells anterior to the morphogenetic furrow (right of the arrowhead) in larvae grown on caffeine(B), whereas posterior to the furrow (left of the arrowhead), caffeine-independent apoptosis is due to the expression of GMR-hid3R (both A and B).



Figure 3-4: Degeneration of $huc95^{E}$ hemizygous imaginal discs in the presence of caffeine. Brains from third instar larvae grown in the presence of caffeine added at day 4 AEL were stained with cleaved caspase-3 and hoescht. Hemizygotes exhibit greater caspase-3 staining at imaginal leg discs while heterozygotes show no staining. DNA stain is added to aid in the visualization of imaginal discs.

	Caspase-3	PH3	DNA	Merge
<i>huc</i> 95 ^E /Df No Caffeine	A	В	С	D
<i>huc</i> 95 ^E / <i>Df</i> Caffeine	E	F	G	H
<i>huc</i> 95 ^E /TM3 No Caffeine		J	К	L
<i>huc</i> 95 ^E /TM3 Caffeine	M	N	0	Ρ

Figure 3-5: Apoptosis and cellular proliferation in eye-antennal discs from $huc95^{E}$ hemizygotes in the presence and absence of caffeine. Third instar larval eye-antennal discs from $huc95^{E}$ hemizygotes raised in the presence (E-F) and absence (A-D) of caffeine were stained with antibodies to caspase-3, PH-3, and hoescht (o visualize apoptosis, mitosis, and DNA respectively). Eye-antennal discs from $huc95^{E}$ heterozygotes in the presence (M-P) and absence (I-L) of caffeine were also stained for a comparison.

	Caspase-3	PH3	DNA	Merge
<i>huc</i> 95 ^E /Df No Caffeine	A	В	C	D
<i>huc</i> 95 ^E / <i>Df</i> Caffeine	E	F	G	Η
<i>huc</i> 95 ^E /TM3 No Caffeine	1	J	К	L
<i>huc</i> 95 ^E /TM3 Caffeine	M	N	0	P

Figure 3-6: Apoptosis and cellular proliferation in wing discs from $huc95^{E}$ hemizygotes in the presence and absence of caffeine. Third instar larval wing imaginal discs from $huc95^{E}$ hemizygotes raised in the presence (E-F) and absence (A-D) of caffeine were stained with antibodies to caspase-3, PH-3, and hoescht (o visualize apoptosis, mitosis, and DNA respectively). Wing discs from $huc95^{E}$ heterozygotes in the presence (M-P) and absence (I-L) of caffeine were also stained for a comparison.



Figure 3-7: Possible model for the caffeine-induced apoptosis in $huc95^{E}$ hemizygotes. Normally, insulin signalling stimulates phosphoinositide 3-kinase (PI3K), which in turn activates protein kinase B (PKB). PKB inhibits apoptosis, thereby allowing cellular proliferation and growth. Inhibition of PI3K by caffeine serves to block activation of PKB, making the cell susceptible to apoptosis. Inhibition of PI3K prevents fusion and allocation of GLUT receptors into the membrane thereby decreasing glucose transport, which can trigger apoptosis. Overexpression of Myo1C in 3T3-L1 adipocyte cells can override PI3K inhibition to allow fusion, which may also be the case with $huc95^{E}$ (if it is myo95E).



Figure 3-8: Mechanism of autophagy. Conjugates of Atg5:Atg12 and Atg8 (LC3 in mammals) is localized to the preautophagosomal structure (PAS). The PAS closes in on itself to form an autophagosome, which is coupled with the loss of Atg5 from the PAS. Fusion of the autophagosome to the lysosome forms an autolysosome, where the membrane and its contents are degraded. (Figure adapted from Rusten *et al.*, 2004)

Chapter 4: Conclusion

As of the writing of this thesis, the identity of $huc95^{E}$ is still unknown. However, out of 23 initial candidates, I have managed to reduce the number to 9, which should make identification easier through molecular-based methods. Out of these candidates, myo95E is of particular interest in that $huc95^{E}$ hemizygotes have a mutation in this gene which obliterates a splicing acceptor site at the junction of intron 8 and exon 9, which is predicted to result in a non-functional protein. Alternative splicing predictions bypassing this mutation by using typical splice sites are predicted to also result in a protein with no function as a myosin, and all of the predicted full-length transcripts utilize this splice site, suggesting that a mutation at this site would have effects on myosin function. Furthermore, RT-PCR analysis confirmed the splicing defect, suggesting that the gene is expressed as well as the splicing mutation. However, a genomic rescue construct containing the full myo95E gene as well as upstream and downstream regions was not able to relieve the caffeine sensitivity of the $huc95^{E}$ mutants. This might be the result of inadequate expression of the construct or could mean that this gene is not a candidate for $huc95^{E}$. If not, several other interesting candidates exist within region one of which is CG6129. This gene is thought to encode a ciliary rootlet component with homology to the Nek2-associated protein C-Nap1, a core centrosomal protein that is responsible for holding the centrioles together during interphase. Nek2 has been implicated in formation of the mitotic spindle and phosphorylation of C-Nap1 by Nek2 at the G₂/M transition is required for centrosomal splitting by displacing C-Nap1 (Rapley et al., 2005; Fry et al., 1998a; Fry *et al.*, 1998b). Furthermore, another related kinase Nek11 is involved in a caffeine-sensitive checkpoint for DNA damage and genotoxic stress (Noguchi et al.,

2002). Since the premise of the screen was to find novel cell cycle checkpoint mutants, this gene is an interesting candidate to pursue and currently efforts to sequence this gene are under way. CG5510 is also an interesting candidate in that it may be involved in secretion and intracellular transport. Other than this, the remaining candidate genes are CG13605, CG13607, CG5463, Kal-1, CG6164, and CG13606 and continued efforts should be able to either eliminate or confirm the status of these genes as candidates for $huc95^{\rm E}$.

 $huc95^{E}$ hemizygotes exhibit pupal lethality in the presence of caffeine and examination of hemizygous pupae reveal no adult structures. In light of this, I have demonstrated that imaginal discs within $huc95^{E}$ third instar hemizygotic larvae exhibit increased apoptosis when exposed to caffeine. This suggests that the caffeine-induced pupal lethality of the $huc95^{E}$ mutants is due to caffeine-induced apoptosis within adultforming tissues, resulting in the inability to complete metamorphosis and might allude to a temporal role for $huc95^{E}$ in development. The role of $huc95^{E}$ in apoptosis within the imaginal discs is unclear and will have to be studied further. Interestingly, $huc95^{E}$ hemizygotes are quite sensitive to caffeine as levels as low as 0.25 mM is enough to compromise their viability (as compared to heterozygous controls which can withstand concentrations of 2 mM caffeine). The role that caffeine plays in this sensitivity is also unclear and is of particular interest in that caffeine is a commonly used reagent in research as well as the most highly consumed drug in the world today.

First and foremost, the identity of $huc95^{E}$ should be determined as such identification could help to elucidate the nature of the caffeine sensitivity displayed by the $huc95^{E}$ mutants. *myo*95E is a promising candidate, containing a mutation that is

predicted to produce a non-functional myosin and an attractive model of how a mutation in this gene could impart caffeine sensitivity has been formulated however, without a suitable rescue to confirm this gene, myo95E only still remains an attractive candidate along with 8 other genes within the region. Currently, efforts are underway to both sequence the remaining genes and to isolate more $huc95^{E}$ alleles through a screen for caffeine sensitive mutants. Hopefully, these efforts will serve to identify the gene responsible once and for all. Furthermore, although the original premise of the screen that had isolated $huc95^{E}$ was to isolate novel cell cycle checkpoint mutants, no work has been done to test whether *huc*95^E has any real checkpoint function. It was determined that huc95E was not sensitive to ionizing radiation, however, this could have been due to redundant checkpoint pathways still present within the *huc*95^E mutant. To address this, the same experiment could be performed but this time, the larvae could be treated with caffeine a day prior to irradiation and eye-antennal or wing imaginal discs could be analyzed for mitotic events. This experiment would test the ability of caffeine to override checkpoint mechanisms (an important assumption in the design of the screen) and if $huc95^{E}$ is involved with a checkpoint response, then perhaps it will be more sensitive to both caffeine treatment and ionizing radiation. Furthermore, the nature of the caffeine sensitivity should be further studied. It was previously determined that caffeine induces apoptosis within the imaginal discs of $huc95^{E}$ hemizygotes however, how or why this occurs still remains a mystery. By determining what pathways are affected by caffeine in these mutants (ie. whether caffeine directly acts on an apoptotic pathway or whether apoptosis is mediated through caffeine's effects on PI3K signalling or DNA repair or checkpoint pathways) could help to elucidate the nature by which $huc95^{E}$ mediates

caffeine sensitivity. It would also be interesting to determine if $huc95^{E}$ is also sensitive to other adenosine analogues (such as purine) and whether these analogues can mediate the same effect as caffeine. Hopefully through this analysis, a better picture will emerge as to why only the imaginal discs seem to be affected and what $huc95^{E}$'s role is in development. Identification and further characterization of $huc95^{E}$ may lead to newer insights into the mechanism of caffeine and its many effects as the world's most popular drug.

Section 4-2: Literature cited

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Appendix

Sequencing Primers: sequences in bold were also used in RT-PCR analysis

Mpk2

		RP-A2	gtgetetacacetegegaet
LP-B1	caagagctgctatcagtcgc	RP-B2	atacggcgacacaaacgaat
LP-C1	atgctctagttgggagtccg	RP-C2	cacgggcacttcattcaaac
LP-D1	taagggtgtcgcatttgctt	RP-D2	gttgcgcaatgagctaaagg
LP-E1	ggttgggcaagtgcagatag	RP-E2	cateteegegetgtacataa
LP-F1	tacggctgcagagtcacata	RP-F2	ccacgagattcggaacttc
LP-G1	atgcaaatcaagttggtcagg	RP-G2	cgacgaggaggtttatcacc
LP-H1	ccatgettateccggtgataa	RP-H2	taccaaatactgcggggact
LP-I1	aacaaccccggcactatgta	RP-I2	cgagttcgttcgttttctgg
LP-J1	gtctcaccttagccacctgg	RP-J2	tcacctcttgggcaggatct
LP-K1	gatgcgtgttaggcattcat	RP-K2	gcaacgccgtgtgtttattt
LP-L1	agcccatataccgtaaaatttgg	RP-L2	gttcgtcggagaatggtagg
LP-M1	cgccggggttatcaaataag	RP-M2	caatgttcccgctatctgag
LP-N1	egactegcaaacatgcagat		888888
Mpk2 Wave 7B	tagacccaaagatcctgcccaagag	Mpk2 Wave 7A	ttgcccgtagacaaatggaaggaat
Mpk2 Wave 6B	cctgagcatacgatggtgggggctt	Mpk2 Wave 6A	catctggtcggtgggctgcatcat
MpK2 Wave 5B	ggtgaatatggtcggtgcctgggaa	Mpk2 Wave 5A	gatgcagcacttgtccgacgaccac
Mpk2 Wave 4B	tcettgacggcaatgtttgagggct	Mpk2 Wave 4A	ccatgcaaagaggacgtaccggga
Mnk2 Wave 3B	tccagcagaccgattacgttctcat	Mpk2 Wave 3A	togatataaatcgaacggaatggga
Mpk2 Wave 2B	cctgtccgtaagctcccgatcccac	Mpk2 Wave 2A	gcategaattatcettcecategca
Mpk2 Wave 1B	toccatocoaacoataattccatoc	Mpk2 Wave 1A	cocatteotototeoettetoateo
I P Mnk2 102	ccatterctaceateeteege	RP Mnk2 201	cagaceteaceaceataegat
LP Mpk2 102	ttaoctootoaatatoottooo	RP Mpk2 207	tcatggccgaactaattaccag
I P Mpk2 103	caagccacggagtatctggt	RP Mpk2 202	gatggacgcagatctgaacaa
LP Mpk2 104	caaacaactacctttaacac	RP Mpk2 203	aatcoaacooaatoooaoat
LP Mpk2 105	tccacgcccaagaatatca	RP Mpk2 204	accentegrantgegutant
LF WIPK2 100	lecaegeeedagaalatea	RP Mpk2 205	caaatttacaatatatagacttta
		101 Mp/2 200	ouuuuuogguuugggouuu
mRnS24			
Intepo24			
L1	tcotaaacaaooccatcaaa	R1	tecatetetagegaacegtta
12	gaccaccatagaggatotot	R2	tccatctctagcgaaccgtta
	EucoBoonnangenBrei	112	iccatere in Begnate Bria
- 20 -			
р38с			
T 1		D 1	
	agegggtgactaaatgeaag		tgecagateeggaagatta
L2	tactcgcgcgttcgaaaaggat	KZ	micgggiaccamcgag
L3	aaggtgcgattttcatcacc	R3	gtatgcgcacctcggctattt
		K4	catectectectetegaat
CG6178			
CG6178-TPI-LPI	gctgtgatcgtatggtggtg	CG6178-TP1-RP1	acaatgtttccgggcagtag
CG6178-TP1-LP2	aattagttcggccatgatgc	CG6178-TP1-KP2	aggetttcaagegaacaete
CG6178-TP1-LP3	tcagatctgcgtccatcaag	CG6178-TPI-RP3	gcaacgccgtgtgtttattt
CG6178-TP1-LP4	atatecgggatetecatte	CG6178-TP1-RP4	caaatgtagcgcattcgtgt
CG6178-TP1-LP5	gatcagaagcgacacacgaa	CG6178-TP1-RP5	cccttcctcaagcaatacc
CG6178-TP1-LP6	aaatttgagcagcaccatcc	CG6178-TP1-RP6	gctggagaacttccaacagg
CG6178-TP1-LP7	ctccccaaaggcgtgtataa	CG6178-TPI-RP7	caacggagaacgagatgaca
CG6178-TP2-LP1	gaacctggactggcaaacat	CG6178-TP2-RP1	cctgtgctggatgagcagta

CG6178-TP2-LP2	gtacggctggcatacatcct	CG6178-TP2-RP2	cgacatggaaatccatcagt
CG6178-TP2-LP3	gacacageetgaetteaega	CG6178-TP2-RP3	agaatgcgacgcagaatctt
CG6178-TP2-LP4	gctcttcctttctgccattg	CG6178-TP2-RP4	agccaatatcgccagtatgc
CG6178-TP2-LP5	ttggttgcatactggcgata	CG6178-TP2-RP5	cacagcaggaccatcaaaga
CG6178-TP2-LP6	ttcaactgaccgagaacgaa	CG6178-TP2-RP6	agcaggttcatttgggtcag
CG6178-TP2-LP7	tgtgacatcgctgaagctct	CG6178-TP2-RP7	ggacaagttaatggcgtgg
myo95E			
TDI I DI MUOUSE	atacaactacotacotacot	TD1_DD1_muo05E	cotataotaoataoaoata
TP1 LP2 Myc05E	tangangetantananganga	TD1 PD2 mv005E	
TP1 I P2 My 095E	actettectttetacetta	TD1 DD2 mv005E	agagetterageteratterage
TP1 + D4 My 095E	atagaagtagaggggg	TD1 DD4 myo05E	tettongoetttongoogo
TP1 LP5 Mvo05E	totacagagagagagagagaga	TP1 PP5 myo05E	actotacactacacttactot
TP1 LP6 Myc05E	totottaattaattaa	TD1 DD6 muo05E	ggicigegiggaeliegiai
TD1 LD7 Myc05E		TD1 DD7 myc05E	
TP1-LF7-My093E		TP1-RP7-IIIy095E	legigaagieaggeigigie
TP2-LP1-My095E	tgggitcantgtgitteca	TP2-RP1-my095E	aagegetgeettttateaga
TP2-LP2-My095E	tetgettttggcacaateet	TP2-RP2-my095E	tegeceteataateegtaae
TP2-LP3-my095E	aagatgacagtctgctggcc	TP2-RP3-my095E	atttggcattcccttcactg
TP2-LP4-my095E	aaagcgatgacgtggatagc	TP2-RP4-my095E	agaaggtcacagccagcatt
TP2-LP5-myo95E	tgtagcacttccaacgatgg	TP2-RP5-myo95E	tatccacgtcatcgcttttg
TP2-LP6-myo95E	ttccttgcgtgatacattcg	TP2-RP6-myo95E	ggccagcagactgtcatctt
TP2-LP7-myo95E	gttacggattatgagggcga	TP2-RP7-myo95E	aggattgtgccaaaagcaga
TP3-LP1-myo95E	gttacggattatgagggcga	TP3-RP1-myo95E	gaaaatccgtcaggcagtgt
TP3-LP2-myo95E	gtgctgggctttttggtaac	TP3-RP2-myo95E	agagcctggaaaaggaggag
TP3-LP3-myo95E	aatgcgagaccattccaatc	TP3-RP3-myo95E	gtaaggttctgggcgatctg
TP3-LP4-myo95E	tgttcgtgctattcgctctg	TP3-RP4-myo95E	gcaagaatacctccgcactt
TP3-LP5-myo95E	aaaattcatgcggtcctcct	TP3-RP5-myo95E	aggaggaccgcatgaatttt
TP3-LP6-myo95E	aagtgcggaggtattcttgc	TP3-RP6-myo95E	acgacaagcagctaccaacc
TP3-LP7-myo95E	ctcctccttttccaggctct	TP3-RP7-myo95E	cttttgcgagttcagcaatg
TP4-LP1-myo95E	etecteetttteeaggetet	TP3-RP8-myo95E	gttaccaaaaagcccagcac
TP4-LP2-myo95E	tgtgggtcagaaagttcacg	TP4-RP1-myo95E	tggcttagcaaatggtttcc
TP4-LP3-myo95E	acagtaccgagtcccaggtg	TP4-RP2-myo95E	gcttgacgcactccttcttc
TP4-LP4-myo95E	agtetetgagecaceaateg	TP4-RP3-myo95E	tttgtggagcgggagaatac
TP4-LP5-myo95E	gtattctcccgctccacaaa	TP4-RP4-myo95E	cgattggtggctcagagact
TP4-LP6-myo95E	gaagaaggagtgcgtcaagc	TP4-RP5-myo95E	catcgttcccgttcttgttt
		TP4-RP6-myo95E	ttaggaggctggcccattat
		TP4-RP7-myo95E	aatccgtcaggcagtgtttc
5'-myo95E-L1	actgaagcgcgaagatttgt	5'-myo95E-R1	gtgctccgtctactgcacaa
5-myo95E-L2	tcgtaaacaaggccatcaaa	5'-myo95E-R2	ccaaggaggtcacagtaccc
5'-myo95E-L3	gacaaacttgcggatgaaca	5'-myo95E-R3	ccagttcgcacatagtctgg
5'-myo95E-L4	cagaaacaattttgcgtcca	5'-myo95E-R4	gtttccggatttcgagacag
5'-myo95E-L5	agttgtcggtgggaaaatgt	5'-myo95E-R5	cgctccagtgctaccgatta
5'-myo95E-L6	agaagcaaagctgcaaaggt	5'-myo95E-R6	ccgttacgtcacgtcaatca
5'-myo95E-L7	tgaggctatcgcagatgatg	5'-myo95E-R7	agacccgccatagaggatgtg
5'-myo95E-L8	ggtgtgcttcgcaatctttc	5'-myo95E-R8	acttcgctgctgttgttttg
-		-	
Df(3R)W6 Analysis

dMyt+ATG LP-CG6178-1 LP-CG6182-1 LP-CG13607-2 LP-CG13605-1 LP-CG13605-1 LP-CG5463-1 LP-CG5510-1 LP-CG13606-1 LP-CHORD-1 LP-CHORD-1	gacggatccatggaaaagcatcatcgcctg gtacggctggcatacatcct tggacgcaaaattgtttctg gtattctcccgctccacaaa tgaggcatcagttcagacg ccgcataatcggaaaaagtg ctgggcattgtggatcttct gcgaagattcagtccgtctc gctggttttgcaccagatt acgacattctgtcggtttcc ttggcttttgctttgc	dMyt-12 RP-CG6178-1 RP-mRpS24-1 RP-CG6182-1 RP-CG13607-2 RP-CG18428-1 RP-CG13605-1 RP-CG6129-1 RP-CG5463-1 RP-CG5510-1 RP-CG13606-1 RP-CHORD-1 PP-Apc2	atagggcggaacacggactct ttatccacgatggggtgttt gagcagctcctccgtgtaac tggcttagcaaatggtttcc tcgtgcatgtgtgtatgtgg ttagctatttgcgacgcttg gctttggcattgggaattta ggccatttgtagcctggtaa cagagcacgcaccactaaaa ccggataatctcatcgtgct acatgtgtacgccatgcact actgggaaaagtccgaggtt
LP-CHORD-1	agaatettgteggeaceate	RP-CHORD-1	actgggaaaagtccgaggtt
LP-Apc2	eteggggaetateceatetea	RP-Apc2	gctccacctgcttctttgac
LP-Tsc1-2	teggttaagggageagetta	RP-Tsc1-2	tgctttccagctgtgatgtc