Raf-mediated cardiac hypertrophy in adult Drosophila

Lin Yu1, Joseph Daniels1, Alex E. Glaser1 and Matthew J. Wolf1,*

SUMMARY

In response to stress and extracellular signals, the heart undergoes a process called cardiac hypertrophy during which cardiomyocytes increase in size. If untreated, cardiac hypertrophy can progress to overt heart failure that causes significant morbidity and mortality. The identification of molecular signals that cause or modify cardiomyopathies is necessary to understand how the normal heart progresses to cardiac hypertrophy and heart failure. Receptor tyrosine kinase (RTK) signaling is essential for normal human cardiac function, and the inhibition of RTKs can cause dilated cardiomyopathies. However, neither investigations of activated RTK signaling pathways nor the characterization of hypertrophic cardiomyopathy in the adult fly heart has been previously described. Therefore, we developed strategies using Drosophila as a model to circumvent some of the complexities associated with mammalian models of cardiovascular disease. Transgenes encoding activated EGFRA\textsuperscript{4887T}, Ras85DV\textsubscript{12} and Ras85DV\textsubscript{12S35}, which preferentially signal to Raf, or constitutively active human or fly Raf caused hypertrophic cardiomyopathy as determined by decreased end diastolic lumen dimensions, abnormal cardiomyocyte fiber morphology and increased heart wall thicknesses. There were no changes in cardiomyocyte cell numbers. Additionally, activated Raf also induced an increase in cardiomyocyte ploidy compared with control hearts. However, preventing increases in cardiomyocyte ploidy using \textit{fuzzy-related} (Fzr) RNAi did not rescue Raf-mediated cardiac hypertrophy, suggesting that Raf-mediated polyploidization is not required for cardiac hypertrophy. Similar to mammals, the cardiac-specific expression of RNAi directed against MEK or ERK rescued Raf-mediated cardiac hypertrophy. However, the cardiac-specific expression of activated ERK\textsuperscript{V23M}, which promotes hyperplasia in non-cardiac tissues, did not cause myocyte hypertrophy. These results suggest that ERK is necessary, but not sufficient, for Raf-mediated cardiac hypertrophy.

INTRODUCTION

Cardiomyopathies are generally associated with a myocyte growth program that leads to an increase in the size of individual muscle cells. Individuals who have cardiac hypertrophy and cardiomyopathies are predisposed to the development of heart failure (Vasan et al., 1997). Heart failure affects 5.7 million individuals in the United States, has an annual economic health care burden in excess of US$34 billion in the United States, is associated with significant morbidity and has a 5-year mortality rate of ~50% despite current pharmacological and device-based therapies (Roger et al., 2010). Furthermore, the development of new pharmaceutical agents to treat heart failure has been disappointing despite an increased understanding of the pathophysiology of cardiomyopathies.

Since the initial descriptions of cardiac hypertrophy in humans, substantial efforts have been directed towards understanding the underlying molecular mechanisms. In response to a variety of stimuli, including RTK-mediated signals, the mammalian heart undergoes morphological changes that contribute to the development of dilated or hypertrophic cardiomyopathies (Heineke and Molkentin, 2006). Dilated cardiomyopathies are characterized by enlargement of the heart chambers, thinning of the heart walls and poor contractility of the myocardium (Braunwald and Bonow, 2012). These changes are manifest as enlargements of chamber dimensions during diastole, when the heart is relaxed, and systole, when the heart is contracted, resulting in systolic dysfunction. In fact, the thinning of the heart walls can result from a process called ‘eccentric hypertrophy’ in which sarcomeres are added in series (Braunwald and Bonow, 2012). Hypertrophic cardiomyopathies are characterized by a ‘thickened but nondilated left ventricle’ (Braunwald and Bonow, 2012). The thickened heart wall in cardiac hypertrophy can occur by the addition of sarcomeres in parallel or by a process whereby the normal architecture of the myocardium becomes disarrayed. As a result, the end-diastolic chamber dimensions are normal or reduced, and systolic function is preserved until overt heart failure develops.

The inhibition of RTKs in the mammalian heart contributes to the development of dilated cardiomyopathies in which the heart chamber becomes enlarged and poorly contractile (Crone et al., 2002). In fact, individuals who receive certain chemotherapy antagonists directed towards RTKs are predisposed to developing dilated cardiomyopathy and heart failure (Chen et al., 2008; Chu et al., 2007; Suter et al., 2007). Conversely, mutations that cause inappropriate activation of RTKs and downstream signaling molecules – such as the small GTP-ase Ras and the serine/threonine-specific protein kinase Raf – are associated with a variety of human syndromes, including Noonan syndrome (Gab and Tartaglia, 2011; Pandit et al., 2007). Moreover, subsets of individuals with Noonan syndrome that have activating mutations in Raf are predisposed to hypertrophic cardiomyopathy (Pandit et al., 2007). Therefore, identifying the signals that cause cardiac hypertrophy can lead to new insights into the pathophysiology of this disease.

Strategies using the fruit fly, \textit{Drosophila melanogaster}, have been developed to facilitate the discovery of genes that cause cardiomyopathies. The adult fly heart is a contractile tube that is a single myocyte layer thick and is composed of pairs of

\textsuperscript{1}Duke University Medical Center, 321 Sands Building, Research Drive, Durham, NC 27710, USA

\*Author for correspondence (wolf0008@mc.duke.edu)

Received 15 November 2012; Accepted 24 March 2013

© 2013. Published by The Company of Biologists Ltd

This is an Open Access article distributed under the terms of the Creative Commons Attribution Non-Commercial Share Alike License (http://creativecommons.org/licenses/by-nc-sa/3.0), which permits unrestricted non-commercial use, distribution and reproduction in any medium provided that the original work is properly cited and all further distributions of the work or adaptation are subject to the same Creative Commons License terms.
DMM

RESEARCH ARTICLE

we identified flies that had enlarged hearts due to mutations in a (ESDs), when the heart is fully relaxed or contracted, respectively, end-diastolic dimensions (EDDs) and end-systolic dimensions mutants causing dilated cardiomyopathies in the fruit fly (Yu et al., syndromes. a model to validate human gene mutations associated with disease signaling pathways, provides unique opportunities to examine how resources available to readily manipulate the fly genome and simplicity of the fly heart, in conjunction with the vast mammals (Bodmer and Venkatesh, 1998; Wolf and Rockman, understanding the normal heart progresses to cardiac hypertrophy and heart failure, the molecular signaling pathways and gene expression programs that together control the growth and differentiation of cardiomyocytes need to be studied in depth.

Results
In this study, the authors provide the first description of cardiac hypertrophy and the signaling pathways controlling its development in adult Drosophila. Using optical coherence tomography (a method that produces images of the heart in intact live flies, similar to echocardiography in humans), confocal microscopy and histological analysis, the authors show that fly cardiomyocytes become enlarged and the volume of cardiac lumen decreases in response to the cardiac-specific expression of signals that emanate from EGFR through Ras, Raf, MEK and ERK. These findings recapitulate aspects of cardiac hypertrophy that are generally observed in mammals. Moreover, cardiac hypertrophy seems to be independent of increases in cardiomyocyte DNA content, an association that has been observed in human cardiac hypertrophy.

Implications and future directions
These results serve as a proof-of-principle that the Drosophila heart can undergo cardiac hypertrophy similar to humans in response to molecular signals. Thus, these findings establish the Drosophila model of cardiac hypertrophy as a platform for the identification of signaling molecules that cause or modify the development of cardiovascular disease. These signaling molecules could potentially provide new therapeutic targets for the treatment of heart failure.

TRANSLATIONAL IMPACT

Clinical issue
Cardiac hypertrophy is a common condition that is triggered by environmental or genetic cues in which the heart muscle enlarges because of an increase in the size of cardiomyocytes. If untreated, cardiac hypertrophy can progress to heart failure and sudden cardiac death. Since the initial descriptions of cardiac hypertrophy in humans, substantial efforts have been directed towards understanding the molecular mechanisms that underlie this condition. Thus, abnormalities in receptor tyrosine kinase (RTK)-mediated signals and downstream signaling molecules are now known to contribute to some forms of cardiac hypertrophy. In addition, specific activating mutations in human Raf, a serine/threonine-specific protein kinase, have been associated with the development of cardiac hypertrophy in subsets of individuals who have Noonan syndrome, a genetic developmental disorder characterized by short stature, specific facial features and congenital heart defects. However, to understand how the normal heart progresses to cardiac hypertrophy and heart failure, the molecular signaling pathways and gene expression programs that together control the growth and differentiation of cardiomyocytes need to be studied in depth.

RESULTS

Activated EGFR or Ras causes cardiac hypertrophy in adult flies
RTKs, including EGFR, are necessary to maintain cardiac function in mammals and flies (Heineke and Molkentin, 2006). Previously, we identified that the inhibition of EGFR caused dilated heart chambers in adult flies (Yu et al., 2010). Because the effects on cardiac function of signaling molecules that are downstream from EGFR have not been evaluated in the adult fly, we examined the effects of EGFR-mediated signaling in the adult fly heart. The tinC-mediated, cardiac-specific expression of constitutively activated EGFR^{A887T} (designated EGFR^{Act}) or the downstream GTPase Ras85DV12 resulted in significantly decreased frequencies of eclosion and poor survival of adult escapers (supplementary material Fig. S1A and data not shown). The embryonic dorsal vessels in tinC>{EGFR^{Act}} heterozygous flies were similar to controls, but the pupal heart in tinC>{EGFR^{Act}} flies in both eclosors and non-eclosors developed similar morphological abnormalities to each other, consistent with the effects on cardiac remodeling during pupal morphogenesis (supplementary material Fig. S1B; Fig. S2). Adult tinC>{EGFR^{Act}} and tinC>{Ras85DV12} heterozygote flies had smaller heart chambers with reduced EDDs compared with controls, as measured by OCT [EDD 95.6±2.9 μm for tinC-Gal4 driver alone (controls) versus 36.4±6.0 μm for tinC>{EGFR^{Act}} or Rhomboid protease. Rhomboid protease performs a crucial step in the processing of a fly ortholog of the EGFR ligand, Spitz, and the expression of soluble forms of Spitz rescued the cardiac abnormalities associated with Rhomboid deficiencies (Yu et al., 2010). Moreover, the cardiac-specific expression of dominant-negative EGFR caused a progressive dilated cardiomyopathy in adult flies, suggesting that EGFR-mediated signals in the Drosophila heart recapitulates abnormalities observed in mammals, including humans.

Dilated heart phenotypes in the fly have been well described; however, the characterization of hypertrophic heart phenotypes in Drosophila has been limited. Using multiple approaches, including OCT, confocal microscopy and histological analyses of heart wall thicknesses, we identify that the activation of EGFR and the downstream molecules Ras and Raf causes a reduction in cardiac lumen sizes at end-diastole and an increase in heart wall thicknesses in adult Drosophila. Furthermore, the changes in heart wall thicknesses are due to myocyte hypertrophy: cardiomyocyte cell number was unchanged. We also show that the cardiac-specific expression of RNAi directed against mitogen-activated protein kinase kinase (MEK) or extracellular signal-regulated kinase (ERK) rescue Raf-mediated cardiac hypertrophy. However, the cardiac-specific activation of ERK does not cause cardiac hypertrophy. This suggests that additional ERK-independent signals emanating from Raf might contribute to hypertrophic cardiomyopathy. In addition, activated Raf induced an increase in cardiomyocyte ploidy compared with that seen in control hearts. However, preventing increases in cardiomyocyte ploidy using fuzzy-related (Fzr) RNAi did not rescue Raf-mediated cardiac hypertrophy, suggesting that Raf-mediated polyploidization is not required for cardiac hypertrophy. Collectively, these results demonstrate that the fly can recapitulate aspects of hypertrophic cardiomyopathy that are seen in mammals and can serve as a platform for the discovery of novel signals that mediate cardiac hypertrophy.

Implications and future directions
These results serve as a proof-of-principle that the Drosophila heart can undergo cardiac hypertrophy similar to humans in response to molecular signals. Thus, these findings establish the Drosophila model of cardiac hypertrophy as a platform for the identification of signaling molecules that cause or modify the development of cardiovascular disease. These signaling molecules could potentially provide new therapeutic targets for the treatment of heart failure.
Cardiac hypertrophy in adult Drosophila

32.3±5.9 μm for tinC>Ras8SDV12, P<0.05; Fig. 1A,B; supplementary material Table S1]. Histological analyses of cardiac chambers from adult tinC>EGFRAct and tinC>Ras8SDV12 heterozygous flies showed heart wall thicknesses that were increased two- to three-times compared with controls (6.7±0.6 μm for controls versus 15.2±0.1 μm for tinC>EGFRAct versus 23.1±1.0 μm for tinC>Ras8SDV12, P<0.05; Fig. 1C,D). To examine the cardiomyocyte architecture in more detail and to avoid obscuration of the heart by the closely attached non-cardiac dorsal diaphragm (also known as the ventral longitudinal muscle), we generated transgenic flies harboring GFP

Fig. 1. The cardiac-specific expression of activated EGFR or Ras8SD causes hypertrophy. (A) OCT M-mode images of the heart lumen from tinC-Gal4 driver control, tinC>EGFRAct and tinC>Ras8SDV12 adult flies. A 1-second marker and 150 μm standard are shown. (B) Summary data for OCT measurements of EDDs for tinC-Gal4 driver control, tinC>EGFRAct and tinC>Ras8SDV12 adult flies. Ten to twelve flies per group. *P<0.05 for indicated transgenic fly line versus control. (C) Histological sections in longitudinal and transverse orientations showing heart wall thicknesses in the first and second abdominal segments (A1/A2) from tinC-Gal4 driver control, tinC>EGFRAct and tinC>Ras8SDV12 adult flies. Arrows indicate heart walls; arrowheads indicate heart wall thicknesses. Scale bar: 10 μm. (D) Summary data for heart wall thicknesses measured from serial transverse histological sections from tinC-Gal4 driver control, tinC>EGFRAct and tinC>Ras8SDV12 adult flies. *P<0.05 for indicated transgenic fly line vs control. Three to six flies per group. (E) Confocal microscopy of tinC-GFP; tinC-Gal4 adult fly heart, tinC>EGFRAct adult fly heart, and tinC>Ras8SDV12 adult fly heart from the A1/A2 abdominal segments. GFP imaging (green) and phalloidin staining of actin (red) show the circumferentially oriented fibers of the adult heart. Ventral longitudinal muscle fibers (also known as the dorsal diaphragm), which do not express tinC-GFP, are also seen by staining with Texas-Red–phalloidin (red) (denoted by arrows). TO-PRO-3 was used for DNA staining (blue). (F) Confocal microscopy with z-stack reconstructions showing the abnormal circumferentially oriented heart myofibers in tinC-GFP; tinC-Gal4 adult fly heart, tinC>EGFRAct adult fly heart, and tinC>Ras8SDV12 adult fly heart from the A1/A2 abdominal segments. GFP imaging (green) and phalloidin staining of actin (red) show the circumferentially oriented fibers of the adult heart. Ventral longitudinal muscle fibers (also known as the dorsal diaphragm), which do not express tinC-GFP, are also seen by staining with Texas-Red–phalloidin (red) (denoted by arrows). TO-PRO-3 was used for DNA staining (blue). (G) Confocal microscopy with z-stack reconstructions showing the abnormal circumferentially oriented heart myofibers in tinC-GFP; tinC-Gal4 adult fly heart, tinC>EGFRAct adult fly heart, and tinC>Ras8SDV12 adult fly heart from the A1/A2 abdominal segments. GFP imaging (green) and phalloidin staining of actin (red) show the circumferentially oriented fibers of the adult heart. Ventral longitudinal muscle fibers (also known as the dorsal diaphragm), which do not express tinC-GFP, are also seen by staining with Texas-Red–phalloidin (red) (denoted by arrows). TO-PRO-3 was used for DNA staining (blue).
Fig. 2. The expression of Ras85D^V12S35, but not Ras85D^V12G37 or Ras85D^V12C40, causes cardiac hypertrophy. Summary data for OCT measurements of EDD (A), ESD (B) and FS (C) for tinC-Gal4 driver control (n=18), tinC>Ras85D^V12S35 (n=11), tinC>Ras85D^V12G37 (n=16) and tinC>Ras85D^V12C40 (n=12) adult flies. *P<0.05 for indicated transgenic fly line vs control. (D) Representative M-mode OCT images for tinC-Gal4 driver control, tinC>Ras85D^V12S35, tinC>Ras85D^V12G37 and tinC>Ras85D^V12C40 adult flies. A 1-second marker and 150 μm standard are shown. (E) Histological sections in transverse orientation showing heart wall thicknesses in the A1/A2 segments from tinC-Gal4 driver control (n=10), tinC>Ras85D^V12S35 (n=11), tinC>Ras85D^V12G37 (n=10) and tinC>Ras85D^V12C40 (n=6) adult flies (top) and summary data of wall thicknesses (bottom). (F) Confocal microscopy with z-stack reconstructions show abnormalities in the heart myofibers in tinC-GFP; tinC>Ras85D^V12S35, tinC-GFP; tinC>Ras85D^V12G37 and tinC-GFP; tinC>Ras85D^V12C40 flies as compared with tinC-GFP; tinC-Gal4 controls. Scale bar: 50 μm. (G) Fluorescent imaging of nuclear-localized RFP in hearts from tinC-RFP^nuc control, tinC-RFP^nuc; tinC>Ras85D^V12S35, tinC>RFP^nuc; tinC>Ras85D^V12G37 and tinC-RFP^nuc; tinC>Ras85D^V12C40 flies. The A2 and A3 abdominal segments are shown.
under the direct control of the cardiac-specific tinC genomic element in the context of the tinC-Gal4 driver (designated tinC-GFP; tinC-Gal4). Hearts from tinC-GFP; tinC-Gal4 flies had circumferentially oriented fibers specifically in cardiomyocytes, and these fibers were readily distinguished from the longitudinal muscle fibers of the dorsal diaphragm (supplementary material Fig. S3). Confocal microscopy and z-stack reconstructions of hearts isolated from tinC-GFP; tinC>Gal4 and tinC-GFP; tinC>Ras85D12 flies showed abnormal cardiac morphology, with myofiber disarray compared with controls (Fig. 1E,F).

Heart wall thicknesses can increase because of an increase in cardiomyocyte number (hyperplasia), an increase in the size of individual cardiomyocytes (hypertrophy) or a combination of both processes. To distinguish among these possibilities, we engineered transgenic flies that harbored GFP and nuclear-localized RFP under the control of tinC (designated tinC-GFP; tinC>RFP nuc). Hearts from tinC-GFP; tinC>RFP nuc/EGFRAct and tinC-GFP; tinC>RFP nuc/Ras85DV12 had the same number of cardiomyocytes as controls, indicating that the increased heart wall thickness was the result of cardiomyocyte hypertrophy, not an increase in cell proliferation (Fig. 1G; supplementary material Fig. S4). Because the activation of EGFR and Ras is known to cause cellular proliferation in other tissues and organs, we validated the effects of the EGFR Act and Ras85D12 transgenes in other tissues and observed proliferative responses in the wing (data not shown). These findings demonstrate that the cardiac-specific activation of EGFR and the downstream signaling component Ras cause cardiac hypertrophy in the fly. Moreover, the tissue context in which EGFR or Ras is activated dictates different cellular responses that lead to proliferation or hypertrophy.

**Ras-mediated cardiac hypertrophy occurs via signals directed to Raf**

Ras is a nodal point in signal transduction and directs the activation of multiple, distinct downstream signal molecules. Therefore, we examined the effects of activated Ras mutants that harbor second site mutations in the effector loop and thereby preferentially direct the activation of Raf, RapGDS or phosphatidylinositol-3-kinase (PI3K) signals via Ras85D12S35, Ras85D12G37 or Ras85D12C40, respectively (Bergmann et al., 1998; White et al., 1995). OCT measurements revealed that flies that were heterozygous for tinC>Ras85D12S35 had smaller EDDs compared with tinC>Ras85D12G37, tinC>Ras85D12C40 or control flies (Fig. 2A,B; supplementary material Table S1). Interestingly, flies that were heterozygous for tinC>Ras85D12C40 had enlarged EDDs compared with controls (EDD 103.9±5.6 μm for tinC>Ras85D12C40; Fig. 2A,B,D; supplementary material Table S1). Confocal microscopy of hearts isolated from tinC-GFP; tinC>Ras85D12S35.
flies showed abnormal cardiac morphology compared with hearts isolated from flies expressing Ras85DV12G37 or Ras85DV12C40, or from controls that contained driver alone (Fig. 2F). Histological analyses demonstrated similar heart wall thicknesses for flies expressing Ras85DV12S35 compared with Ras85DV12G37 or Ras85DV12C40 or with controls (Fig. 2E; supplementary material Table S1). These findings were consistent with observations that, although Ras85DV12S35 preferentially signals to Raf, the intensity of Ras85DV12S35-mediated signals is diminished compared with those of Ras85DV12 (White et al., 1995). Additionally, hearts from tinC-GFP; tinC>RFPnuc/Ras85DV12S35 flies and tinC-GFP; tinC>RFPnuc/Ras85DV12C40 flies had the same number of cardiomyocytes, indicating that each Ras transgene did not cause cardiomyocyte proliferation (Fig. 2G).

**Activated fly or human Raf causes cardiac hypertrophy and an increase in cardiomyocyte ploidy in adult flies**

Next, we examined the effects of cardiac-specific expression of fly and human Raf proteins in the adult fly heart. Fly and human Raf constructs that had deletions of the amino acids from 2 to 431 for fly and from 2 to 334 for human Raf (designated fly RafAct and human RafAct) have been shown to possess constitutive activity that is independent of Ras (Brand and Perrimon, 1994). Hearts from flies that were heterozygous for tinC-fly RafAct or tinC>human RafAct had smaller EDDs, cardiomyocyte myofiber disarray and increased heart wall thicknesses compared with controls (Figs 3, 4; supplementary material Table S1). Conversely, hearts from flies that were heterozygous for tinC-RafRNAi lines had increased EDSs, decreased fractional shortening (FS) and thinner heart walls compared with controls (Fig. 3B,C; supplementary material Table S1). Additionally, hearts rates, cardiac cycle durations, systolic and diastolic intervals, and arrhythmia indices were similar between control and RafAct hearts (supplementary material Fig. S5).

Hearts from tinC-GFP; tinC>RFPnuc/fly RafAct and tinC-GFP; tinC>RFPnuc/human RafAct had the same number of cardiomyocytes (Fig. 5A). However, the nuclear RFP signal appeared intense in the hearts expressing RafAct (Fig. 5A). Because RFP expression was non-quantitative, we performed additional quantitative analyses of cardiomyocyte DNA content using established methods (Fox et al., 2010). Hearts that expressed RafAct had an increase in cardiomyocyte ploidy [mean ploidy (C value) was 33 for tinC-GFP; tinC>human RafAct flies versus 22 for tinC-GFP; tinC-Gal4 controls, P<0.05], suggesting that the expression of activated human Raf increased endoreplication (Fig. 5B).

To examine whether the observed hypertrophy is largely, or solely, caused by the increase in ploidy of the cells, we used RNAi directed against Fzr to artificially decrease polyploidization in the presence of activated Raf. Fzr promotes cell cycle progression during Drosophila endocycling in multiple tissues, and postmitotic salivary gland cells fail to enter endoreduplication cycles in Fzr-deficient flies (Sigrist and Lehner, 1997; Zielke et al., 2008). Therefore, we examined the cardiac-specific expression of Fzr RNAi in the context of RafAct. The cardiac-specific expression of Fzr RNAi prevented Raf-mediated increases in cardiomyocyte ploidy [mean ploidy (C value) was 40 for tinC-GFP; tinC>human RafAct flies versus 17 for tinC-GFP; tinC>RafAct + Fzr RNAi flies, P<0.05; Fig. 5C]. Interestingly, heart wall thicknesses were similar between tinC-GFP; tinC>human RafAct and tinC-GFP; tinC>human RafAct + Fzr RNAi (Fig. 5D), suggesting that Raf-mediated increases in polyploidy were not required for cardiac hypertrophy.

Because Raf is known to activate MEK, we examined the effects of MEK knockdown in adult fly hearts. Hearts from tinC>MEKRNAi flies had slightly increased EDDs, significantly increased EDSs, and

---

**Fig. 4. Raf-mediated changes in cardiomyocyte morphology cause concentric hypertrophy in adult Drosophila.** Confocal microscopy with z-stack reconstruction of the A1/A2 segments of the adult Drosophila heart from tinC-GFP; tinC-Gal4 (A,C,E,G) and tinC-GFP; tinC>human RafAct (B,D,F,H) flies, showing disarrayed fibers and increased thickness in single cardiomyocytes in the latter. Arrows denote the fiber orientations in cardiomyocytes and the asterisks denote the heart lumen. Scale bars: 10 μm. Schematic interpretation of the confocal micrographs showing fly cardiomyocytes that are normal (I) or that have concentric hypertrophy (J).
Cardiac hypertrophy in adult Drosophila

thinner heart walls compared with controls (Fig. 6A-D; Fig. 7; supplementary material Table S1). Flies that had cardiac-specific expression of activated Raf in the context of MEKRNAi (tinC>MEKRNAi, tinC>human RafAct) had cardiac chamber dimensions, heart wall thicknesses and cardiomyocyte myofibrillar architecture that were similar to tinC>MEKRNAi (Fig. 6A-D; Fig. 7; supplementary material Table S1). Five independent transgenic UAS-MEKRNAi lines were examined, and four lines rescued the Raf-mediated cardiac abnormalities (supplementary material Table S1 and data not shown), suggesting that Raf-dependent signals to MEK are necessary for Raf-mediated cardiac hypertrophy.

ERK is necessary, but not sufficient, for the development of cardiac hypertrophy in adult Drosophila

ERK is one of the downstream effector molecules through which Raf mediates cellular signals (Heineke and Molkentin, 2006). Activating mutations in rolled, the fly ortholog of ERK, were originally identified from fly screens of EGFR mutants and subsequently shown to result from a mutation of aspartic acid to asparagine at amino acid 334 (Brunner et al., 1994). The aspartate amino acid at position 334 in fly ERK is conserved across species, including humans, and ERKDD34N was shown to be a dominant mutation in the catalytic domain of the kinase (Oellers and Hafen, 1996). Therefore, we generated transgenic constructs harboring UAS-ERKDD34N and validated the activity of the transgene in the fly eye using a GMR-Gal4 driver or in the fly wing using a dpp-Gal4 driver. GMR>ERKDD34N had a rough eye phenotype consistent with the initially characterized fly mutant and dpp>ERKDD34N had additional veins and clonal expansion of wing epithelium, indicating that the ERKDD34N transgenes encoded functionally active ERK (Fig. 6E,F). Interestingly, the cardiac chamber dimensions and heart wall thicknesses in tinC>ERKDD34N were similar to controls (Fig. 6A; Fig. 7A; supplementary material Table S1). Thus, the cardiac-specific expression of activated ERK did not phenocopy the cardiac hypertrophy under conditions of EGFR, Ras or Raf activation. Next, we examined the effects of genetically ablating ERK in the context of activated Raf. tinC>ERKRNAi flies had cardiac parameters that were similar to controls but the heart walls were thinner and had normal-appearing cardiomyocyte myofibrillar structure (Fig. 6A-D; Fig. 7; supplementary material Table S1). Flies with cardiac-specific expression of activated ERK in the context of ERKRNAi (tinC>ERKRNAi, tinC>human RafAct) had cardiac chamber dimensions, heart wall thicknesses and cardiomyocyte myofibrillar architecture that were similar to tinC>ERKRNAi, suggesting that ERK is epistatic to Raf. Thus, the activation of ERK did not cause cardiac hypertrophy. However, the inhibition of ERK prevented Raf-mediated cardiac hypertrophy, suggesting that ERK is necessary but not sufficient (Fig. 8).

Fig. 5. Raf-mediated cardiac hypertrophy is independent of increases in cardiomyocyte ploidy. (A) Fluorescent imaging of nuclear-localized RFP in hearts from tinC>RFPnuc controls and tinC>RFPnuc/fly RafAct. The A2 and A3 abdominal segments are shown. (B) The distribution of cardiomyocyte ploidy (C value) from tinC-GFP, tinC>human RafAct (n=106 nuclei) (black bars) and tinC-GFP, tinC-Gal4 control (n=61 nuclei) hearts (white bars) expressed as the percentage of nuclei of total for each genotype. The mean ploidy C value is shown for each group. *P<0.05 for tinC-GFP, tinC>human RafAct versus tinC-GFP, tinC-Gal4 controls. (C) The distribution of cardiomyocyte ploidy (C value) from tinC-GFP, tinC>human RafAct (n=50 nuclei) (black bars), tinC-GFP, tinC>human RafAct + Fzr RNAi (n=52 nuclei) (white bars) and tinC-GFP, tinC>human RafAct + Fzr RNAi (n=50 nuclei) (gray bars) hearts expressed as the percentage of nuclei of total for each genotype. The mean ploidy C value is shown for each group. (D) Summary data of wall thicknesses from tinC-GFP, tinC>human RafAct (n=6), tinC-GFP, tinC>human RafAct + Fzr RNAi (n=6) and tinC-GFP, tinC>human RafAct + Fzr RNAi (n=5) adult flies. *P<0.05 for tinC>human RafAct + Fzr RNAi versus tinC-GFP, tinC>human RafAct + Fzr RNAi. NS, non-significant.
DISCUSSION

The inhibition of EGF ligand or EGFR by dominant-negative EGFR molecules results in enlarged cardiac chambers in adult flies (Yu et al., 2010). Our current studies show that the cardiac-specific expression of activated EGFR, Ras or Raf (human or fly) causes decreased heart chamber lumens. These results raise the question: how does the lumen of a heart composed of pairs of single myocytes undergo reduction or enlargement (i.e. dilation)? One proposed explanation is that molecular signals that drive the addition of sarcomeres added in series produce an eccentric hypertrophy and resultant enlarged heart lumen. Conversely, signals that promote either the addition of sarcomeres in parallel or myofiber disarray produce enlarged myocytes and resultant concentric hypertrophy in the fly. This explanation is consistent with a model proposed by Molkentin’s group that showed that ERKs regulate the balance between eccentric and concentric cardiac growth in mammals (Kehat et al., 2011). Transgenic mice that had genetically ablated ERK1 and ERK2 in the heart (Erk1−/−, Erk2fl/flαMHC-Cre mice) developed an eccentric hypertrophy, and isolated cardiomyocytes from these mice displayed elongation. Our

Fig. 6. The inhibition of MEK or ERK prevents Raf-mediated cardiac dysfunction. Summary data for OCT measurements of EDD (A), ESD (B) and FS (C) for tinC-Gal4 driver controls (n=18), tinC>human RafAct (n=18), tinC>ERKD334N (n=12), tinC>human RafAct with MEK RNAi (n=15), tinC>ERK RNAi (n=15) and tinC>human RafAct with ERK RNAi (n=15). *P<0.05 for indicated transgenic fly line vs control. (D) Representative M-mode OCT images for tinC-Gal4 driver controls, tinC>human RafAct, tinC>MEK RNAi, tinC>human RafAct with MEK RNAi, tinC>ERK RNAi, and tinC>human RafAct with ERK RNAi. A 1-second marker and 150 μm standard are shown. (E) Expression of ERKD334N in the fly eye using a GMR-Gal4 driver causes a rough eye phenotype (arrowhead). (F) Expression of ERKD334N in the wing using a dpp-Gal4 driver causes clonal expansion of wing epithelium (arrows).
results show that the inhibition of ERK or MEK causes a thinner cardiomyocyte and in some cases an enlargement in EDDs, consistent with findings in mammalian models.

The expression of activated EGFR, Ras or Raf promotes cell proliferation in the eye or wing (Baker and Rubin, 1989; Brunner et al., 1994; Halfon et al., 2000; Karim and Rubin, 1998; Lu et al., 1994). The activation of these pathway components in the fly heart caused cardiac hypertrophy. We observed myocyte polyploidy in control hearts and increases in myocyte ploidy in the RafAct hearts that is suggestive of incomplete endocycling, repetitive rounds of genome replication. An increase in cell ploidy is achieved through endoreplication, in which genomic DNA content increases without cellular division (Lee et al., 2009). Endoreplication is a common occurrence among species and, in the fly, some cells can have up to 2048 copies of the euchromatic genome (Edgar and Orr-Weaver, 2001). This process has been described as an effective strategy for cell growth, often found in differentiated cells that are large or have high metabolic activity.

Prior studies have shown changes in cardiomyocyte DNA content, ploidy level and nuclear number in mammalian hearts across multiple species, including humans (Adler et al., 1996). Post-mortem examinations of human hearts demonstrated that the degree of polyploidy closely correlated with myocardial hypertrophy (Adler and Friedburg, 1986; Sandritter and Scomazzoni, 1964). Recently, endoreduplication has been observed in mouse cardiomyocytes after cardiac injury (Hesse et al., 2012). Augmented endoreduplication, and an increase in ploidy, might represent a mechanism by which cells can adapt to high metabolic demands (Edgar and Orr-Weaver, 2001). Therefore, growth conditions or stimuli that promote cardiac hypertrophy might cause the myocyte to adapt by increasing ploidy, thereby providing more copies of essential genes required to respond to these cues (Ahuja et al., 2007).

In our studies, Fzr RNAi prevented Raf-mediated increases in cardiomyocyte ploidy. Interestingly, the cardiac-specific expression of Fzr RNAi did not significantly change the degree of Raf-mediated cardiac hypertrophy. Therefore, the cardiac hypertrophy observed in the context of activated Raf does not require increased polyploidization. One interpretation of these findings is that the signals emanating from Raf bifurcate towards two distinct pathways: one that drives DNA replication and one that promotes cardiac hypertrophy.

The results further support the concept that the cellular context in which signaling molecules are expressed defines the cell growth response, namely hyperplasia or hypertrophy. During fly development, signals from EGFR are required for the specification and diversification of embryonic muscle progenitors, including cardiac cells (Baylies et al., 1998; Bodmer and Venkatesh, 1998; Carmena et al., 1995; Zaffran and Frasch, 2002). Somatic muscles and the cardiac cells develop from specialized progenitors (Carmena et al., 1995; Carmena et al., 1998; Ward and Skeath, 2000). Each progenitor cell divides asymmetrically and produces two founder cells that specify individual muscle cell fates and give rise to multinucleate myofibers (Rushton et al., 1995).
has shown that hyperactive EGFR, using early mesodermal GAL4 drivers including twist-GAL4 or 24B-GAL4, generates supernumerary mesodermal founder cells and causes a duplication of dorsal acute muscle 1 (DA1), a larval body wall muscle (Buff et al., 1998). Moreover, Wingless and the TGFβ family member Decapentaplegic, two members of the signals from the Wnt family, prepare the mesoderm and render cells competent to respond to Ras-MAPK activation (Halfon et al., 2000). Thus, the timing of EGFR and Ras signals during development and the integration of cellular cues dictates the response of myocytes, including the cardiomyocytes.

We used the cardiac driver tinC-Gal4, which is expressed later in mesodermal development: this might explain why we did not observe the increase in cardiomyocytes that has been observed with drivers that are expressed earlier in mesodermal development (Lo and Frasch, 2001; Zaffran et al., 2006). Our findings suggest that adult Drosophila cardiomyocytes lose the ability to proliferate when EGFR signals are activated after cardiomyocyte differentiation, similar to observations in mammals. Whether this lack of myocyte proliferation is specific to EGFR, Ras and Raf or is more generalizable to other stimuli remains to be investigated.

In mice, the indirect activation of ERKS via upstream signaling molecules has been shown to cause cardiac hypertrophy. Transgenic knock-in mice expressing RafL613V, a mutation that is associated with Noonan syndrome, develop cardiac hypertrophy. Transgenic RNAi Project (TRiP) at Harvard Medical School (http://www.flyrnai.org/). All fly stocks were maintained on standard yeast protein media at room temperature (Ashburner et al., 2000). TheVinna Drosophila RNAi Center (VDRC) RNAi stock directed against Fzr (w^{1118}, P{GD9960}v2550) was kindly provided by Hermann Steller (Bergmann et al., 1998). The described (Wolf et al., 2006; Yu et al., 2010). p{LIAS-Ras85D^V12C40} and p{LIAS-ERKD334N} were engineered by site-directed mutagenesis from cDNA from w^{1118}, and transgenic insertion lines were obtained by standard protocols (Brand and Perrimon, 1993). p{tinC-GFP} ; p{UAS-RFP^NIDC}, p{tinC-Gal4} were generated using p{tinC-GFP} ; p{tinC-Gal4} and p{UAS-RFP^NIDC} transgenic stocks. Transgenic flies harboring p{LIAS-Ras85D^V12C40} and p{LIAS-Ras85D^V12G7} were kindly provided by Don Fox (Duke University, Durham, NC). All other fly stocks were obtained from the Bloomingston Stock Center or the Transgenic RNAi Project (TRiP) at Harvard Medical School (http://www.flyrnai.org/). All fly stocks were maintained on standard yeast protein media at room temperature (Ashburner et al., 2005).

**OCT measurement of cardiac function in adult Drosophila**

Cardiac function in adult Drosophila was measured using a custom-built OCT microscopy system (Bioptigen Inc., Durham, NC) as previously described (Wolf et al., 2006; Yu et al., 2010). Adult female Drosophila between 3 and 10 days post-eclosion were briefly subjected to CO₂, placed on a soft gel support, and allowed to fully awaken based on body movement. All flies were imaged in B-modes.
in the longitudinal orientation to identify the cardiac chamber in the A1 segment and then in the transverse orientation to center the heart chamber. Multiple 3-second OCT M-modes were recorded for each fly. M-mode OCT images were processed using ImageJ software and referenced to a 150-μm standard. After M-modes were acquired, the flies were re-examined in the transverse B-mode orientation to assure consistent measurements from the heart chamber. End-diastolic dimensions (EDDs), end-systolic dimensions (ESDs) and heart rate were determined from three consecutive heart beats. Fractional shortening (FS) was calculated as (EDD–ESD)/EDD×100.

**Histological analysis**

Fly heart wall thicknesses were measured as previously described (Yu et al., 2010). Briefly, adult female flies of 5-7 days age after eclosion were collected and fixed in Telly's fixation buffer (60% ethanol, 3.33% formalin, 4% glacial acetic acid) for at least 1 week at 4°C. Previously, we determined that cardiac chamber sizes in flies from this fixation step were similar to the EDD in awake, adult flies as assessed using OCT. Adult *tinC-EGFR^{64c} and tinC-Ras85D^{12f}* flies were collected at 2-3 days after eclosion because they had impaired survival.

Specimens were dehydrated in ethanol through sequential gradients. Then, the samples were washed twice with xylenes before immersion in liquid paraffin. After solidification, paraffin blocks were sectioned serially at 8 μm thickness in longitudinal or transverse orientation. Sections were rehydrated and stained with hematoxylin and eosin. Established criteria were used to control for the position of the heart chamber among different flies that were evaluated (Yu et al., 2010). Measurements were made in three serial 8-μm sections. Sections were analyzed using a Leica DM2500 microscope equipped with a Leica DFC310FX digital camera. Wall thickness was calculated by measuring the cardiac chamber wall width along the mid-dorsal, mid-ventral, left lateral and right lateral wall in three serial sections to obtain the mean ± s.e.m.

**Evaluation of adult cardiac morphology**

Adult *Drosophila* corresponding to the F1 offspring of *p{tinC-GFP} ; p{tinC-Gal4}* stocks crossed to specific *p{UAS-transgenes}* or *w^{1118}* (controls) were collected at 2-3 days age, post-eclosion, to examine adult cardiac morphology. Flies were briefly anesthetized by administration of CO₂, the head and thorax were removed and the abdomen was placed in artificial hemolymph buffer [108 mM Na⁺, 5 mM K⁺, 2 mM Ca²⁺, 8 mM MgCl₂, 1 mM NaH₂PO₄, 4 mM NaHCO₃, 10 mM sucrose, 5 mM trehalose, and 5 mM HEPES (pH 7.1)]. An incision was made along the ventral aspect of the abdomen and the internal abdominal organs were gently removed. The surrounding fat and tissue were removed using a pulled glass capillary pipette. Hemolymph buffer that contained 10 mM EGTA was then added to relax the cardiac muscle as described by Alayari et al. (Alayari et al., 2009). Next, samples were fixed in 4% paraformaldehyde for 20 minutes at room temperature prior to staining with a primary anti-GFP-antibody (1:500; Invitrogen, Inc.) and secondary antibody conjugated to Alexa Fluor 488 (1:500; Invitrogen, Inc.) for detection of cardiomyocytes, with phalloidin–Texas-Red (1:1000; Invitrogen, Inc.) for actin staining, and with TO-PRO-3 (1:10,000; Invitrogen, Inc.) for DNA staining. The stained heart preparations were visualized under a Zeiss LSM510 confocal microscope and 0.4 μm z-stack images were analyzed.

For evaluation of cardiac morphology during pupal stages, *Drosophila* corresponding to the F1 offspring of *p{tinC-GFP} ; p{tinC-Gal4}* stocks crossed to specific *p{UAS-transgenes}* or *w^{1118}* (controls) were collected between the P6 and P13 stages according to staging described by Bainbridge and Bownes (Bainbridge and Bownes, 1981). Pupal hearts were directly visualized using a Leica M165FC fluorescent stereomicroscope. For embryo staining, pericardin was detected using EC11 anti-Pericardin-5. The EC11 anti-Pericardin-5 developed by D. Gratecos was obtained from the Developmental Studies Hybridoma Bank (DSHB), developed under the auspices of the NICHD and maintained by The University of Iowa, Department of Biology, Iowa City, IA.

**Evaluation of adult cardiomyocyte ploidy**

Ploidy was determined by adapting methods previously described (Fox et al., 2010). Adult *Drosophila* corresponding to the F1 offspring of *p{tinC-GFP} ; p{tinC-Gal4}* stocks crossed to specific *p{UAS-transgenes}* or *w^{1118}* (controls) were collected at 2-3 days age, post-eclosion, and hearts were dissected in artificial hemolymph buffer as described above. Testes from adult male flies were isolated as internal controls because this tissue is haploid. Dissected hearts were then removed from the cuticles. Hearts and testes were placed on the same coverslip for all subsequent steps. Samples were fixed in 4% paraformaldehyde at room temperature for 20 minutes, then washed three times (1 minute per wash) in TBT buffer (10 mM Tris-HCl pH 8.0, 250 mM NaCl, 0.1% BSA, 0.1% Triton X-100) prior to staining nuclei with TO-PRO-3 (1:5000 dilution in TBT) for 20 minutes at room temperature in the dark. Samples were then washed three times (1 minute per wash) in TBT prior to adding VECTASHIELD, applying a positively charged glass slide (Sigma) and squashing the slide for 10 seconds. The samples were then analyzed by obtaining z-stacks of TO-PRO-3-stained nuclei using confocal microscopy. Cardiomyocytes were identified by live GFP expression. Identical emission intensities were used for each slide to quantify TO-PRO-3 staining of myocyte and testis nuclei. The intensity of each myocyte or testis nuclei was quantified using ImageJ, and myocyte nuclei TO-PRO-3 expression was normalized to testis nuclei TO-PRO-3 expression for each slide to calculate ploidy. Ploidy was expressed as the C-value, where a C-value of 1 refers to the amount of DNA contained within a haploid nucleus.

**Heart rate analyses**

Adult *Drosophila* corresponding to the F1 offspring of *p{tinC-GFP} ; p{tinC-Gal4}* stocks crossed to specific *p{UAS-human Raf^{64c} or w^{1118}}* (controls) were collected at 2-3 days age, post-eclosion, and hearts were dissected in artificial hemolymph buffer as described above. Heart movements were recorded using a Leica GPF stereo microscope equipped with an Andor iXon X3 EMCCD camera at a rate of 100 frames per second for 10 seconds and processed using Solaris software (Andor, Inc.). Each recording was processed using Excel software to identify the local minima and maxima and corresponding time-stamped image frame. The systolic interval (SI), diastolic interval (DI), RR (peak-to-peak interval), CC (cardiac cycle length), fraction of cardiac cycle in systole (SI/CC), fraction of cardiac cycle in diastole (DI/CC), and the arrhythmia index (AI)
are shown. AI was calculated as the standard deviation of the RR interval normalized to the median of the RR interval as previously described (Fink et al., 2009).

Statistical analysis
Comparisons of EDD chamber dimensions were determined by either a Student's t-test for two samples or an analysis of variances (ANOVA) with corrections for multiple comparisons when necessary. GraphPad Prism (GraphPad Software, Inc.) and Microsoft Excel statistical software were used for all analyses.

ACKNOWLEDGEMENTS
We thank the TRIP at Harvard Medical School (NIH/NIGMS R01-GM84947) for providing transgenic RNAi fly stocks and/or plasmid vectors used in this study. We thank Dr Van Bennett for use of his confocal microscope. We are grateful to Don Fox and Howard Rockman for helpful discussions.

COMPETING INTERESTS
The authors declare that they do not have any competing or financial interests.

AUTHOR CONTRIBUTIONS
L.Y., J.D., A.E.G. and M.J.W. conceived, designed and performed the experiments, analyzed the data and wrote the paper.

FUNDING
This work was supported by the National Institutes of Health (NIH) (R01HL116581) and the American Heart Association Mid-Atlantic Beginning Grant-In-Aid (128GA1191007) to M.J.W., and the American Heart Association Mid-Atlantic Post-Doctoral Award (10POST320036) to L.Y.

SUPPLEMENTARY MATERIAL
Supplementary material for this article is available at http://dmm.biologists.org/lookup/suppl/doi:10.1242/dmm.011361/-/DC1

REFERENCES


Suter, T. M., Procter, M., van Veldhuisen, D. J., Muscholl, M., Bergh, J.,
Oncol. 25, 3859-3865.
ventricular dilatation and the risk of congestive heart failure in people without
cells and their progenitors in the Drosophila embryo. Development 127, 4959-
4969.
White, M. A., Nicolette, C., Minden, A., Polverino, A., Van Aelst, L., Karin, M. and
Wigler, M. H. (1995). Multiple Ras functions can contribute to mammalian cell
transformation. Cell 80, 533-541.
Med. 22, 55-61.

(2006). Drosophila as a model for the identification of genes causing adult human
Wu, X., Simpson, J., Hong, J. H., Kim, K. H., Thavarajah, N. K., Backx, P. H., Neel, B.
phenotypes in a mouse model of Noonan syndrome associated with the
457-469.
Cardioblast-intrinsic Tinman activity controls proper diversification and
differentiation of myocardial cells in Drosophila. Development 133, 4073-4083.
Zielke, N., Querings, S., Rottig, C., Lehner, C. and Sprenger, F. (2008). The anaphase-
promoting complex/cyclosome (APC/C) is required for rereplication control in