MEF2 Is an In Vivo Immune-Metabolic Switch

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SUMMARY

Infections disturb metabolic homeostasis in many contexts, but the underlying connections are not completely understood. To address this, we use paired genetic and computational screens in Drosophila to identify transcriptional regulators of immunity and pathology and their associated target genes and physiologies. We show that Mef2 is required in the fat body for anabolic function and the immune response. Using genetic and biochemical approaches, we find that MEF2 is phosphorylated at a conserved site in healthy flies and promotes expression of lipogenic and glycogenic enzymes. Upon infection, this phosphorylation is lost, and the activity of MEF2 changes—MEF2 now associates with the TATA binding protein to bind a distinct TATA box sequence and promote antimicrobial peptide expression. The loss of phosphorylated MEF2 contributes to loss of anabolic enzyme expression in Gram-negative bacterial infection. MEF2 is thus a critical transcriptional switch in the adult fat body between metabolism and immunity.

INTRODUCTION

Metabolic regulation is tightly and ubiquitously linked with immune responses and inflammatory signaling (Hotamisligil, 2006). Prolonged or excessive immune activation can drive metabolic disruption and cause wasting of fatty and lean tissues. This effect is seen in many human infections; it is particularly prominent in Gram-negative sepsis and in persistent bacterial infections such as tuberculosis (Schwenk and Macallan, 2000; Tappy and Chiofò, 2007). Because of the many etiologies behind infection-induced cachexia, numerous molecular mechanisms have been proposed to underlie this condition, with signals including lipid mediators and cytokines and transcription factors including FOXO, NF-κB, AP-1, Stats, and nuclear receptors acting singly or in combination (Tracey and Cerami, 1994; Vallier and Hotamisligil, 2010; Van den Bergh, 2002). It has been a major challenge to the field to experimentally link any one of these molecular mechanisms to observed metabolic dysfunction following infection in vivo.

Chronic or acute infection disrupts systemic metabolism in Drosophila as well as in vertebrates. We have previously shown that infection with Mycobacterium marinum causes metabolic disruptions in Drosophila resulting, in part, from a systemic loss of AKT activity (Dionne et al., 2006). This promotes pathological FOXO activation and an inability to produce new metabolic stores. Activation of the Toll pathway in the fly fat body, either genetically or by infection, is sufficient to partially phenocopy this effect, both at the level of AKT activity and metabolic storage (DiAngelo et al., 2009). Listeria infection causes similar metabolic pathology (Chambers et al., 2012). The function of the link between immune activation and loss of anabolic signaling activity is unclear, especially because FOXO is able to activate antimicrobial peptide expression but is not required for resistance to infection (Becker et al., 2010). How other anabolic or catabolic pathways are altered by infection in flies is also unknown, though it is clear that there are other common regulators of these processes, at least at the level of the whole organism (Rynes et al., 2012).

We used a paired screening strategy to identify the pathways and transcriptional networks that regulate host physiology in vivo in response to infection. One screen used computational analysis of gene expression after different immune challenges to identify coregulated genes and their predicted transcriptional...
regulators. The other screen involved testing mycobacterial infection susceptibility of flies carrying targeted RNAi knock-downs of transcription factors and signaling intermediates. These screens converged on MEF2, a pleiotropic transcription factor originally characterized as a key factor in muscle development (Bour et al., 1995; Lilly et al., 1995; Molkentin et al., 1995). Our data reveal that MEF2 regulates immune and metabolic activities, as depletion of MEF2 in the fat body causes dramatic failures of systemic anabolism and immune function. This results from reduced expression of key metabolic enzymes and antimicrobial peptides, respectively. The choice between immune and metabolic target genes is dictated by phosphorylation of MEF2 at a conserved site. In healthy animals, MEF2 is phosphorylated at T20 and promotes expression of its metabolic targets, whereas infection results in T20 dephosphorylation and association with the TATA-binding protein (TBP) at a distinct TATA sequence of immune targets.

**RESULTS**

**Computational and Functional Screens Reveal Mef2 as a Regulator of Infection Susceptibility**

To identify new factors in the response to infection, we performed two screens. In the first, we analyzed a large microarray data set reflecting whole-fly gene expression at multiple times after several different infections (for details, see Figure 1 and supporting material) (De Gregorio et al., 2001, 2002; Dionne et al., 2006). This data set included wild-type animals as well as loss-of-function mutants in the imd and Toll pathways, the two primary microbe-detection systems in the fly. These two pathways respond to different microbes and share no molecular components but regulate overlapping sets of target genes because each culminates in activation of a distinct NF-κB family member (Lemaitre and Hoffmann, 2007). We used fuzzy c-means clustering to find clusters of genes coregulated across biological conditions and then identified transcription factor binding sites overrepresented in the vicinity of the genes within each cluster (Figure 1 and Tables S1 and S2 available online). These binding sites represent potential regulators of the associated genes.

This type of computational analysis affords direct identification of transcriptional effectors, their targets, and regulated physiologies simultaneously without prior knowledge. This is reflected in the fact that many of our coregulated groups of genes correspond to clear biological functions. In keeping with our previous observation of metabolic dysregulation in Mycobacterium marinum-infected flies, metabolic functions were overrepresented in 6 of the 30 clusters (Figure S1), suggesting that the shared transcriptional regulators of these clusters may have important metabolic roles.
To test in vivo function of predicted regulators, we performed a secondary functional screen, testing time to death after *M. marinum* infection in flies with ubiquitous or fat-body-specific RNAi knockdown of 29 transcription factors or signals associated with specific factors predicted to directly regulate gene clusters after infection. Five genes gave clear survival phenotypes; of these, *Mef2* was particularly intriguing, being associated both with immune-response genes (cluster P) and with metabolic activities (cluster X). We further analyzed *Mef2* as a potential nodal point between immune activation and metabolic disruption.

**Mef2 Regulates Susceptibility to Infection**

The fat body, a homolog of mammalian liver and adipose tissue, is both the primary source of the inducible humoral immune response and the site of metabolic stores in the fly. Because strong loss-of-function mutants in *Mef2* are embryonic lethal (Bour et al., 1995; Lilly et al., 1995), we tested animals with *Mef2* RNAi knockdown driven by c564, a driver strongly expressed in the fat body (Hrdlicka et al., 2002). Flies carrying fat body *Mef2* RNAi exhibited a significant reduction in survival time after a high-dose *M. marinum* infection compared with driver-only controls (Figure 2A). This survival effect was strengthened at lower bacterial doses.

To determine whether MEF2 is generally required for survival after infection, we assayed survival of fat body *Mef2* knockdowns after other infections. In addition to *M. marinum*, these animals were compromised in response to a second intracellular pathogen, *Listeria monocytogenes* (Figure 2B). Moreover, unlike wild-type animals, they were killed by infection with either *Enterococcus cloacae* or *Candida albicans* (Figures 2C and 2D). Defense against these infections is mediated specifically by the *imd* and TOLL pathways, respectively (Gottar et al., 2006). Sensitivity of flies lacking fat body *Mef2* to *E. cloacae*, in particular, was similar to that of *imd* pathway mutants (Figure 2A) (Gottar et al., 2006; Hedengren et al., 1999). This could be explained if these flies lacked the fat body, but we found no visible reduction in fat body mass or decrease in c564-driven GFP expression (Figure S2A). Other fat body drivers recapitulated the susceptibility to *E. cloacae* (Figure S2D), and the susceptibility resulting from *Mef2* knockdown was rescued by coexpression of wild-type *Mef2* (Figure S2E).

We then assayed bacterial numbers in animals infected with *E. cloacae* or *M. marinum*. In each case, more bacteria were present in *Mef2* knockdown animals (Figures 2E and 2F). The dramatic increase in *E. cloacae* number suggests that the effect on this infection may be fully accounted for by defects in bactericidal activity. By contrast, the modest difference observed in *M. marinum* number suggests that, in a more complex, chronic infection, MEF2 is also important in tolerance of pathology.

**Mef2 Is Required for Normal Fat and Glycogen Storage**

As mentioned earlier, our data set included many genes encoding metabolic activities. Cluster X, a predicted MEF2 target
cluster, contained several key metabolic enzymes, including acetyl-CoA carboxylase (ACC), diacylglycerol acyltransferase (mdy), and fat body hexasse kinase (Hex-C). In addition to being repressed by M. marinum infection, cluster X was rapidly and transiently downregulated 3–6 hr after septic injury (infection with a mixture of nonpathogenic Gram-positive and Gram-negative bacteria), with most genes returning to baseline within 24 hr (Figure 1B). In addition to genes in cluster X, genes encoding key anabolic enzymes in clusters Y and AA, as well as unclustered genes, were repressed after either M. marinum infection or septic injury. These observations led us to examine the effects of MEF2 on the expression of cluster X genes—and other anabolic enzymes showing similar expression—in the absence of infection.

MEF2 knockout reduced expression of several key enzymes in fatty acid and triglyceride synthesis, including ACC, mdy, fatty acid synthase (FASN/CG3523), ATP citrate lyase (ATPCL), glycerol-3-phosphate dehydrogenase (Gpdh), glycerol kinase (Gk), and the mitochondrial citrate transporter (sea) (Figure 3A). A similar pattern was apparent for the enzymes of fat body glucose uptake and glycogen synthesis, including Hex-C, Trett-1 (one of two GLUT family members highly expressed in fat body [Chintapalli et al., 2007]), phosphoglucose mutase (Pgm), UDP-glucose pyrophosphorylase (UGP), glycogen synthase (GlyS/CG6904), and the glycogen branching enzyme (AGBE) (Figure 3A, black bars). The expression of almost all these genes was rescued by coexpression of wild-type Mef2 (Figure 3A, gray bars); for a few targets, this rescue was partial, likely because the RNAi targeted the rescue transgene as well as the endogenous locus. Genes that were incompletely rescued or that showed paradoxical repression in the rescue-containing animals (FASN and Hex-C) could also be repressed by inducible expression of a Mef2-engrailed repressor fusion (Blanchard et al., 2010), confirming their dependence on Mef2 for expression (Figure S3A). However, the fat body was intact and continued to express glycogen phosphorylase (GlyP), GFP driven by c564, and other fat-body-enriched genes, indicating that Mef2 is not required for development, survival, or overall identity of this tissue (Figures S3A and S2).

In keeping with reduced expression of enzymes of glycogen synthesis (Hex-C, Pgm, UGP, GlyS, and AGBE) and lipogenesis (sea, ATPCL, ACC, FASN, Gpdh, Gk, and mdy), unchallenged Mef2 knockout flies were almost entirely devoid of triglyceride and glycogen (Figures 3B, 3C, S3B, and S3C). These animals contained normal levels of free glucose and trehalose, suggesting that, even without glycogen or triglyceride stores, they can regulate circulating sugar (Figure 3B). These metabolic phenotypes could be seen, though more weakly, with a second fat body driver (Figures S3D and S3E; data not shown). Other fat body drivers may give weaker phenotypes because they knock down Mef2 less efficiently or because c564 also knocks down Mef2 in other tissues (for example, the gut, which is lipogenic and parts of which also exhibit c564-driven expression). As would be expected, Mef2 knockout animals died more rapidly than wild-type flies when starved (Figure S3F). However, they were not generally stress sensitive, exhibiting normal survival under either hyperoxia or heat stress (Figures S3G–S3J).

These data suggest that the statistical association of predicted Mef2 sites with genes encoding key metabolic activities reflects a requirement for Mef2 in normal expression of these genes and thus in normal metabolic function. Although MEF2 may regulate its metabolic targets indirectly, we favor direct regulation, as 12/13 of the identified metabolic target genes have high-quality MEF2 binding sites within their 5’ or 3’ flanking

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**Figure 3. Metabolic Effects of Mef2 Knockdown**

(A) Expression of genes encoding enzymes of triglyceride and fatty acid synthesis, glucose uptake, and glycogen synthesis in control (wt; c564/+), Mef2 knockout (Mef2 knockdown; UAS-Mef2-IR/+), and rescued (Mef2 rescue; UAS-Mef2-IR/+; UAS-Mef2-IR/+; Mef2 knockdown) flies assayed by qRT-PCR. mRNA levels are normalized to the control genotype (wt; c564+/+) and expressed as log2. Values are mean ± SD. Significance levels: *p < 0.01 and **p < 0.001 by heteroscedastic unpaired two-tailed t test.

(B) Total triglyceride, glycogen, and free glucose measured by colorimetric enzymatic assay in Mef2 knockdown flies compared with driver-only controls.

(C) Neutral lipids stained by oil red O in driver-only control and Mef2 knockdown fat body. Red, oil red O; blue, DAPI. Scale bars, 20 μm. See also Figure S3 and Table S3.
regions or in introns (Table S3). The severe metabolic defect in Mef2 knockdown animals likely contributes to their short lifespan after mycobacterial infection by decreasing their ability to withstand infection-induced wasting.

**Mef2 Knockdown Impairs Humoral Immune Responses**

Although Mef2 has not previously been described as an immune regulator in *Drosophila*, our computational analysis also associated Mef2 sites with cluster P, which contains many genes known to be targeted by NF-κB-like factors upon activation of the *imd* or *Toll* pathways, including the antimicrobial peptides Defensin, Metchnikowin, Drosocin, and Attacin A. These genes are typically strongly induced within 3 hr of septic injury.

To test whether MEF2 regulates these genes, we examined antimicrobial peptide response to septic injury in fat body Mef2 knockdown flies. We again infected flies with a mixture of the *imd* agonist *Escherichia coli* and the *Toll* agonist *Micrococcus luteus*. These are strong agonists of the respective pathways but do not rapidly kill flies in which detection pathways are mutated, permitting us to assay gene expression at late times after infection. Fat body Mef2 knockdown flies exhibited dramatically reduced induction of *AttA*, *CecA1*, *Def*, *Dpt*, *Dro*, *Im4*, and *Mtk* in response to this mixed infection (Figure 4, compare black bars to white bars). Again, this could be rescued by coexpression of wild-type *Mef2* (Figure 4, gray bars), and again, the few genes that were not completely rescued also exhibited reduced induction in flies carrying an inducible *Mef2-EnR* transgene (Figure S4A). This block was heterogeneous; at least one antimicrobial peptide, Listericin, was entirely unaffected (Figure 4).

We observed similar results in animals that received single infections with either bacterium (Figure S4B). That both *Toll* and *imd* target genes were affected in response to an infection activating both pathways indicates that the role of MEF2 is not specific to either pathway.

**MEF2 and TBP Physically Interact upon Infection to Bind AMP TATA Boxes**

To clarify the requirement for Mef2 in expression of its immune targets, we examined the requirement for individual MEF2 sites near antimicrobial peptides. We cloned putative regulatory regions (~1.5 kb to the ATG) from *Drosocin* and *Metchnikowin* upstream of enhanced GFP (eGFP). This region included NF-κB, DEAF-1, and GATA sites previously shown to regulate *Mtk* expression (Reed et al., 2008; Senger et al., 2004) as well as multiple putative MEF2 sites. In each case, a predicted Mef2 site overlapped the TATA box (*Dro* site 3 and *Mtk* site 2) (Figures 5A and S5). Individual putative MEF2 binding sites were mutated in these constructs, keeping the core TATA sequences intact; each Mef2 mutant TATA sequence replicated a naturally occurring, functional TATA box not predicted to bind MEF2. We initially analyzed reporter activity of wild-type and mutant constructs in S2* cells treated with lipopolysaccharide (LPS) (commercial LPS contains sufficient peptidoglycan to potently activate the *imd* pathway [Kaneko et al., 2004] (Figure S5). Both Dro-GFP and Mtk-GFP were strongly induced by LPS, as expected. Distal Mef2 sites were important for *Dro* reporter activity, but both Mtk and Dro reporters required the MEF2-TATA box for normal induction (Figure S5). To test in vivo responses to infection, we made transgenic flies carrying the same eight GFP reporters, each inserted into the AttP2 site. Uninfected, these animals exhibited weak or no GFP fluorescence, but *E. coli* injection induced strong fat body GFP activity in flies carrying the wild-type reporters.
The requirement for MEF2 sites in vivo was similar to that in vitro; in particular, the MEF2-TATA site was required in each case (Figure 5A).

We then tested whether MEF2 and TBP interacted directly by immunoprecipitating endogenous MEF2 from whole-fly lysates. In adult flies infected with the \textit{imd} agonist \textit{E. cloacae} or a mixture of \textit{M. luteus} and \textit{E. coli}, TBP coimmunoprecipitated with MEF2 (Figure 5B and data not shown). When stimulated by \textit{E. cloacae}, this association required the \textit{imd}-activated JNKKK Tak1 but was unimpaired in \textit{Dif}; \textit{Rel} double mutants, suggesting a requirement for JNK signaling, but not NF-κB, in MEF2-TBP complex formation after Gram-negative infection (Figure 5B).

We next asked whether the MEF2-TBP complex could bind the putative MEF2-TATA site. The \textit{Dro} MEF2-TATA site (TCTATATAAAGC) was a near-perfect match to the ideal MEF2A site as defined by SELEX (KCTAWWWWTAGM) (Pollock and Treisman, 1991), so we focused on the more divergent \textit{Mtk} MEF2-TATA site (GCTATAAAAGC). This site eliminates one of the central A/T nucleotides in the MEF2 consensus, changing the spacing between the two putative half-sites. When assayed by EMSA, the \textit{Mtk} core promoter was bound by nuclear extracts from S2R+ cells transfected with \textit{Mef2}-HA, causing two distinct complexes to form: complex 1, at lower apparent molecular weight, was strong and consistently present, whereas the amount of complex 2 varied from experiment to experiment, though it was always less abundant than complex 1 (Figure 5C).

Mutation of the putatively MEF2-binding C and G nucleotides flanking the TATA box eliminated both complexes. Complex 1 was also eliminated by preincubation of extracts with antibodies to either HA or TBP, indicating that this complex contained MEF2 and TBP together bound to the \textit{Mtk} TATA sequence.

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To test the generality of these observations, we examined the promoters of 30 antimicrobial peptides (AMPs) and other immune-inducible small peptides (Table S4). Of these 30 genes, 27 had a TATA box, defined as TATAWAAGM, is identical to the Mtk MEF2-TATA box (Figure 5D and Table S4). 17/27 AMP TATAs match this consensus; 3/27 instead match the Dro site, CTATA TAAAGC. The remaining seven TATA boxes include at least one nonconsensus nucleotide in the C...AGM flanks. The Mtk and Dro-like sites were found on peptides from many structural families, implying that this is not an ancestral sequence maintained by lack of counterselection. Conversely, the 13 metabolic genes shown in Figure 3A had a total of 27 TSSs; of these, 5 had TATA boxes, none of which matched the extended consensus associated with antimicrobial peptides.

We then examined roughly 15,000 unique promoters drawn from four databases of precisely mapped Drosophila TSSs (Hoskins et al., 2011; Ni et al., 2010; Rach et al., 2009; Schmid et al., 2006). This identified 890 TATA-containing promoters, with TATA boxes defined by the same criteria used for antimicrobial peptides (Table S5). The median TATA box position was 31 nucleotides upstream of the TSS (25–75 percentile range, 30–33 nucleotides) (Figure S5B). This set of TATA boxes contained 13 sequences that had been in our “training set” of AMPs and AMP-like peptides; these were not removed—and the other AMP TATAs were not added to this set—to avoid functionally biasing the represented genes. AMP promoters were more likely to contain a TATA box than those in the unbiased set (27/30 TATA-containing AMP promoters versus 890/15,000 TATA-containing promoters in the unbiased analysis). Unlike AMP TATA boxes, the 890 unbiased TATA boxes showed little nucleotide preference outside the core TATAWA, except a preference for A or T in the immediate 3’ position (Figure S5D).

If the nucleotides flanking the core TATAWA motif permit regulation of TATA usage via MEF2-TBP interaction, then the identities of these nucleotides should be correlated—the presence of a 5’ C should predict the presence of a 3’ AGM and vice versa. We examined the relationships among flanking nucleotide positions within the 890 unbiased TATA boxes. In each case, TATA boxes that fit the Mtk-like TATA consensus at one flanking nucleotide were more likely to match this consensus at other positions as well. For example, in the initial set, 27.1% of TATA boxes contained a C directly 5’ of the TATAWA core. Fixing the nucleotide 3’ of the TATAWA as A increased this representation to 34.2%, fixing the two 3’ nucleotides as AG gave a further increase to 38.3%, and fixing the three 3’ nucleotides as AGM increased the representation to 41.4% (Figure 5E).

We extracted from our set of TATA-containing promoters those matching the Mtk-like or Dro-like consensus. This gave 88 genes (Table S5). In addition to antimicrobial peptides, this list contained other infection-regulated genes and genes involved in responses to a variety of stresses. It also contained larval cuticular proteins, genes involved in sensory development or function, and 13 different uncharacterized small proteins with predicted secretory signals. The Gene Ontology terms “humoral immune response,” “immune response,” “defense response,” and “antimicrobial humoral response” were overrepresented among genes with MEF2-TATA sequences as compared with the full set of 890 TATA-containing genes (Bonferroni-corrected p < 0.01).

These data together indicate that infection drives formation of a MEF2-TBP complex that can bind the TATA box of many antimicrobial peptides and is required for normal transcriptional induction of these genes. That the flanking nucleotide positions do not vary independently and the MEF2-TBP TATA box is overrepresented on immune genes both suggest that the MEF2-TBP TATA box represents a previously uncharacterized discrete gene regulatory element.

MEF2 Is Phosphorylated In Vivo at T20 to Regulate Association with TBP

The mechanisms by which the infection-inducible MEF2-TBP complex might be regulated and its potential function in MEF2 regulation of metabolic targets were unclear. We returned to our microarray data to directly compare expression of the metabolic and immune genes we had identified as Mef2 regulated. After bacterial infection, these genes were strikingly counterregulated; when the antimicrobial genes were induced, expression of the metabolic genes was lost. This was especially pronounced 3 and 6 hr after septic injury and in late-stage mycobacterial infection (Figure 6A). This observation suggested that MEF2 modification might underlie a choice between immune and metabolic targets.

MEF2 activity can be regulated by several mechanisms, including phosphorylation by p38 MAP kinase, a calcineurin-regulated acetylation/sumoylation switch, and an indirect mechanism involving phosphorylation-regulated recruitment of class II histone deacetylases (Berdeaux et al., 2007; Han et al., 1997; Shalizi et al., 2006; Zhang et al., 2002). In agreement with previous work (Chen et al., 2010; Dijkers and O’Farrell, 2007), neither p38 nor calcineurin significantly altered immune competence in adult flies (data not shown).

MEF2 phosphorylation at T20, within the DNA-binding domain, can promote binding of MEF2 to a consensus MEF2-dimer binding sequence that does not contain a central TATAWA (Wang et al., 2005). The T20 region is conserved in Drosophila, C. elegans, and all mouse and human MEF2 homologs, and it matches a crude consensus site for AKT and p70 S6 kinase (Figure 6B). Using a phosphomotif antibody (Moritz et al., 2010), we found that MEF2 T20 phosphorylation was detectable in extracts from whole-adult Drosophila and was reduced in flies that had received weak or strong immune challenges (PBS injection or mixed infection with M. luteus and E. coli, respectively) (Figure 6C).

To test the relevance of this phosphorylation in transcriptional complex formation, we generated flies that inducibly expressed HA-tagged wild-type MEF2, nonphosphorylatable MEF2 (T20A), or phosphomimetic MEF2 (T20E) specifically in the fat body. One week after eclosion, these flies were shifted from 18° to 29° to drive Mef2*HA expression. Because all transgenes were inserted into the same site, Mef2* constructs were expressed at similar levels (Figure S6A).

We tested whether T20 directly regulated the association of MEF2 with TBP by immunoprecipitating fat-body-expressed
Figure 6. T20 Phosphorylation Regulates MEF2-TBP Association

(A) Counterregulation of Mef2 targets after infection. Data are from microarrays discussed in text. Expression levels are normalized to uninfected controls. Values are mean ± SD.

(B) The Mef2 T20 region is conserved in flies, humans, and C. elegans and matches an AKT/p70 S6 kinase consensus site.

(C) Endogenous MEF2 immunoprecipitated from whole flies is phosphorylated at T20; this phosphorylation is reduced by PBS injection or by infection with mixed M. luteus and E. coli. Top indicates immunoprecipitated MEF2 probed with anti-MEF2 antibody; below indicates the same blot probed with anti-phospho-RXRXXS/T* antibody; bottom indicates quantification of phospho-T20-MEF2 from the blot shown, normalized to total MEF2 and to the level in unmolested flies.

(D) Immunoprecipitation (IP) of inducible fat-body-expressed wild-type, T20A, or T20E MEF2.HA from flies either uninfected or infected 3 hr previously with mixed E. coli and M. luteus, probed with anti-HA (top) or for coimmunoprecipitated endogenous TBP (bottom). Genotype: w1118/Y; c564/+; UAS-Mef2*.HA/tub-Gal80ts.

(E) T20 can be phosphorylated by p70 S6K in vitro. Purified GST or GST-MEF2tide fusions (wild-type or T20A) were incubated with 32P-g-ATP with or without recombinant kinases and resolved by SDS-PAGE. Top indicates 32P autoradiogram; bottom indicates Coomassie blue stain.

(F) p70 S6K Thr398 phosphorylation is reduced by infection. Top indicates representative western on whole-fly lysates probed with the antibodies indicated; bottom indicates quantification of phospho-S6K normalized to tubulin from three independent experiments. Values are mean ± SD. Infections: Ec = E. coli (Gram-negative IMD agonist); Ml = M. luteus (Gram-positive TOLL agonist). Uninjected animals were anaesthetized alongside injected animals but were otherwise unmolested.

See also Figure S6.
expressing wild-type model, we measured expression of immune and metabolic regulators for antimicrobial peptide expression. To test this, we wished to identify a kinase or kinases responsible for MEF2 T20 phosphorylation in the Drosophila fat body. Others have observed MEF2C T20 phosphorylation by protein kinase A (PKA) and AKT (Wang et al., 2005). As PKA activation inhibits glycosylation and promotes lipolysis and as AKT activity is reduced by infection in Drosophila, we initially focused on AKT as a likely T20 kinase. However, we were unable to observe phosphorylation of this site by AKT in vitro (Figure 6E). Like AKT, p70 S6 kinase (S6K) is broadly anabolic in its effects and is activated by nutrient signals. S6K shares with AKT its preference for substrates with arginine at −5 and −3 (Alessi et al., 1996). T20 was efficiently phosphorylated by recombinant S6K in vitro (Figure 6E), and systemic S6K activation (measured by phosphorylation at T398, corresponding to T389 in human S6K) was reduced in flies that had received infections with Toll or imd agonists (Figure 6F). A similar effect was seen during the last 3 days of life in flies infected with M. marinum (Figure 6G). S6K is thus a plausible in vivo MEF2 T20 kinase and may be responsible for promoting expression of the enzymes of lipogenesis and glycogenesis in healthy, well-fed animals.

### T20 Regulates the Ability of MEF2 to Promote Distinct Target Genes and Physiologies

Together, these data suggested that loss of MEF2 T20 phosphorylation might cause the loss of anabolic transcripts seen after infection and that loss of T20 phosphorylation might be permissive for antimicrobial peptide expression. To test this model, we measured expression of immune and metabolic MEF2 targets in uninfected and infected animals inducibly expressing wild-type Mef2, Mef2T20E, Mef2T20A, or driver-only controls. In most cases, overexpression of wild-type MEF2 had small effects on either the loss of metabolic transcription or the induction of antimicrobial peptides after infection when compared to the effects of infection in driver-only controls (Figures 7A and S7). Expression of nonphosphorylatable MEF2 enhanced expression of antimicrobial peptides (by as much as 6-fold, in the case of Defensin) after either M. luteus or E. cloacae infection but had little effect on most metabolic genes. Conversely, expression of phosphomimetic MEF2 had little general effect on antimicrobial peptides and did not counteract the loss of most anabolic transcripts after M. luteus infection but rescued levels of anabolic transcripts after E. cloacae infection.

Together, these data indicate that T20 phosphorylation can regulate the association of MEF2 with TBP, switching MEF2 between metabolic and immune functions. The loss of metabolic gene expression driven by imd pathway activation (typically caused by Gram-negative bacterial infection) stems, at least in part, from the loss of T20-phosphorylated MEF2.

### DISCUSSION

Here, we identify Mef2 as a factor critical for energy storage and the inducible immune response in the Drosophila fat body. Many infection-induced antimicrobial peptides depend on Mef2 for normal expression. In consequence, flies lacking Mef2 activity in the fat body are severely immunocompromised against a variety of infections. Mef2 sites are also associated with genes encoding key enzymes of anabolism, and Mef2 is required for normal expression of these genes; consequently, flies lacking Mef2 function in the fat body exhibit striking reductions in the total levels of triglyceride and glycogen. These two groups of target genes are counterregulated during infection; the anabolic targets of Mef2 are reduced in expression when antimicrobial peptides are induced. We show that fat body Mef2 can exist in two states with distinct physiological activities. In uninfected animals, Mef2 is phosphorylated at T20 and can promote the expression of its metabolic targets. In infected animals, T20 is dephosphorylated, and Mef2 associates with the TATA-binding protein to bind a compound MEF2-TATA sequence found in the core promoters of antimicrobial peptides. The loss of T20-phosphorylated Mef2 promotes the loss of anabolic transcripts in flies with Gram-negative bacterial infection. These data, taken together, suggest that the central role of MEF2 in promoting fat body anabolism and immune activity reflects a switch between distinct transcriptional states regulated, at least in part, by differential affinity for TBP determined by T20 phosphorylation (Figure 7B).

The signaling mechanisms regulating T20 phosphorylation and MEF2-TBP association are clearly of critical importance. The ability of p70 S6K to phosphorylate this residue is congruent with the ability of S6K to enhance anabolism and repress catabolism in response to nutrient signals (Laplante and Sabatini, 2012). However, others have shown T20 phosphorylation by PKA, suggesting that T20 phosphorylation may be regulated by more than one pathway in vivo (Wang et al., 2005). The role of TAK1 may be similarly complex. TAK1 is required for formation of the MEF2-TBP complex upon Gram-negative infection, but this effect may be indirect. For example, reduced S6K phosphorylation after infection may result from insulin resistance driven by TAK1 via JNK (Chen et al., 2002; Hirosumi et al., 2002). TAK1-dependent JNK activation is required for normal AMP induction in vivo (Delaney et al., 2006), but it remains possible that some novel pathway is the critical connection between TAK1 and MEF2-TBP complex formation.

In mammals, in addition to hematopoietic roles, Mef2c regulates B cell proliferation upon antigen stimulation (Khiem et al., 2008; Wilker et al., 2008), and Mef2d regulates IL2 and IL10 in T cells (Liopeta et al., 2009; Pan et al., 2004). The possibility that Mef2 family proteins might be important direct activators of innate responses has not previously been examined. We show that Mef2 is a core transcriptional component of the innate immune response of the adult fly. Equally, vertebrate Mef2 family proteins are critical regulators of muscle metabolism, activated by physical activity to promote expression of PGC-1α and the glucose transporter Glut4 (Handschin et al., 2003; Thai et al., 1998; Wu et al., 2001). Glut4 regulation by Mef2 is known in adipose tissue as well as in muscle (Thai et al.,
It is an intriguing possibility that MEF2 is as important a regulator of adipose metabolism in vertebrates as we show it to be in flies.

**Figure 7. T20 Phosphorylation Regulates Mef2 Target Choice in Infected Flies**

(A) Effects of inducible fat body expression of wild-type, T20A, or T20E Mef2 assayed by qRT-PCR. Genotypes are as in Figure 6D. Mef2 expression was activated beginning 16 hr before infection. Left indicates the effect of infection in driver-only controls; right indicates changes due to misexpression of Mef2 mutants. Overall significance of changes is assayed by Wilcoxon signed-rank test. All measurements are the mean of three biological replicates.

(B) Model. The position of Tak1 is unresolved: though shown upstream of S6K inactivation, it could also promote MEF2-TBP interaction in parallel with S6K inactivation.

See also Figure S7.

Infection-induced metabolic disruption leading to cachexia is present in vertebrates as well as in insects, most notoriously in Gram-negative sepsis and persistent bacterial infections such as peritonitis.
as tuberculosis (Schwenk and Macallan, 2000; Tappy and Chiovéro, 2007). Our data suggest that wasting seen after infection may be due, in part, to the requirement for MEF2 to serve different transcriptional functions in different conditions; the MEF2 immune-metabolic transcriptional switch may be a mechanistic constraint that forces the fly into metabolic pathophysiology in contexts of persistent immune activation. Alternatively, the loss of MEF2-driven anabolic transcripts due to infection may be productive, either by altering systemic energy usage or by increasing the production or release of one or more antimicrobial metabolites. Recent work has highlighted a distinction between “resistance” type immune mechanisms, in which the host attempts to eradicate an invading organism, and “tolerance” type mechanisms, in which the host response is oriented toward reducing the damage done by infection (Schneider and Ayres, 2008). The distinct metabolic and immune requirements for MEF2, combined with the obligation on the part of the host to raise some measure of resistance to systemic infection, may limit the achievable level of tolerance in persistent infections.

EXPERIMENTAL PROCEDURES

Fly Culture, Stocks, and Infections
Flies were maintained at 25°C, 60% relative humidity, and on food containing 10% w/v Brewer’s yeast, 8% fructose, 2% polenta, and 0.8% agar. Adult males were collected soon after eclosion and transferred to fresh food to age prior to treatment. All phenotypes shown were observed with VDRC Mef2-IR lines 15549 and 15550, which gave essentially identical results. 5- to 10-day-old adult male flies were injected with 50 nl of bacterial suspension or vehicle as described (Dionne et al., 2003). For survival, dead flies were counted at 24 hr intervals or more frequently. Survival analyses were repeated at least three times, always showing qualitatively similar results. Minimum cohort size was 20 flies.

Antimicrobial peptide reporter expression was assayed in live flies using a Leica M205 FA. Photographs were taken and processed using identical settings.

Microarrays
Experimental aspects of array analysis have been described (De Gregorio et al., 2001, 2002; Dionne et al., 2006). Data were reanalyzed using the BioConductor suite in R.

Clustering of Probes, Binding-Site Predictions, and Functional Predictions
We analyzed only probes significantly regulated (fold change > 1.5, p < 0.01) in end-stage *M. marinum* samples (Dionne et al., 2006). Probes were clustered by multiple rounds of fuzzy c-means clustering using MFuzz (Futschik and Carlisle, 2005).

For each cluster, extended gene regions (gene sequence including introns and UTRs, plus 2,000 bases flanking sequence) were analyzed with CLOVER (Firth et al., 2004), using Drosophila chr 3R as background sequence.

Binding sites were tested against 20 false clusters of genes (genes selected randomly from those represented on the Affymetrix Drosophila Genome 1 array and treated the same as true clusters). Sites predicted to regulate more than two false clusters were eliminated from consideration.

Functional predictions were from Generic GO Term Finder (http://go.princeton.edu/cgi-bin/GOTermFinder).

Biochemistry
EMSA, immunoprecipitation, kinase assays, and western blots were performed according to standard protocols as described in the Extended Experimental Procedures.

Quantitative Real-Time RT-PCR
Performed as previously described (Clark et al., 2011; Dionne et al., 2006). All data shown were the result of three to six biological replicates. Oligonucleotides are listed in Table S6.

ACCESSION NUMBERS
The ArrayExpress accession number for the *M. marinum* microarray data reported in this paper is E-MEXP-2510.

SUPPLEMENTAL INFORMATION
Supplemental Information includes Extended Experimental Procedures, seven figures, and six tables and can be found with this article online at http://dx.doi.org/10.1016/j.cell.2013.09.007.

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