Regulation of BMP Signaling by O-GlcNAcylation

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Graphical Abstract

Normal glucose and Sxc

Extracellular space

Cytoplasm

Low glucose or no Sxc

Normal glucose and Sxc

Low glucose or no Sxc
Summary

Precise regulation of signal transduction is critical throughout organismal life, both for embryonic development and for adult homeostasis. To ensure proper spatio-temporal signal transduction, Bone Morphogenetic Protein (BMP) signaling pathways, like all other signaling pathways, are regulated by both agonists and antagonists. Here, we report identification of a previously unrecognized method of signal antagonism for Dpp (Decapentaplegic), a *Drosophila* BMP family member. We demonstrate that the BMP type I receptor Saxophone (Sax) functions as a Dpp receptor in the *Drosophila* embryonic epidermis, but that its activity is normally inhibited by the O-linked glycosyltransferase Super sex combs (Sxc). In wild-type embryos, inhibition of Saxophone (Sax) activity in the epidermis marks the BMP type I receptor Thickveins (Tkv) as the sole conduit for Dpp. In contrast, in sxc mutants, the Dpp signal is transduced by both Tkv and Sax, and elevated Dpp signaling induces errors in embryonic development that lead to embryonic death. We also demonstrate that Sax is the O-glycosylated target of Sxc and that O-glycosylation of Sax can be modulated by dietary sugar. Together, these findings link fertility to nutritive environment and point to Sax (activin receptor-like kinase [Alk] 1/2) signaling as the nutrient-sensitive branch of BMP signaling.

Keywords

Dpp, Tkv, Sax, Alk, Sxc, OGT, GlcNAc, FOP, dorsal closure
INTRODUCTION

Bone Morphogenetic Proteins (BMPs) are an evolutionarily important group of signaling molecules conserved in vertebrates and invertebrates, with evidence suggesting a common ancestor at least 600 million years ago (Padgett et al., 1993). BMPs were first identified and named for their ability to induce bone development (Urist, 1965). Since their discovery, BMPs have been shown to play roles in conserved embryonic developmental and adult homeostatic processes by regulating cell lineage commitment, differentiation, proliferation, apoptosis, and morphogenesis in various cell types throughout the body (Katagiri and Watabe, 2016). These are all processes with a mechanistically undefined, but emerging relationship to O-GlcNAc (Sun et al., 2016). Decapentaplegic (Dpp) is the *Drosophila* orthologue of mammalian BMPs 2 and 4, themselves members of the BMP branch of the Transforming Growth Factor-β (TGF-β) superfamily (Padgett et al., 1987). Disruptions of BMP signaling components exhibit shared loss-of-function developmental phenotypes in metazoans (Moulton and Letsou, 2016; Reiter et al., 2001). Flies, in particular, have provided a powerful platform for discerning the biological activities and the biochemical properties of BMPs and their intracellular signal transducers.

BMPs (including Dpp) are diffusible ligands that form signaling gradients. Upon binding to a receptor complex containing two (homomeric) type I and two (homomeric) type II serine/threonine kinase receptors, the BMP signal is transduced intracellularly via phosphorylation of transcriptional regulators. Both BMP receptor types (I and II) have an extracellular ligand binding domain, a single-pass transmembrane domain, and an intracellular kinase domain (ten Dijke and Hill, 2004). A type I receptor-specific domain, the glycine-serine (GS) domain, is additionally required for full kinase activity (Franzen et al., 1993). When complexed, the BMP type II receptor, a constitutively active serine/threonine kinase, transphosphorylates the GS domain of its BMP type I receptor partner. In its turn, the activated BMP type I receptor phosphorylates and activates a receptor Smad (R-Smad), which subsequently enters the nucleus and binds to BMP-responsive
targets to activate and enhance transcription (Massague, 1998). Measures of R-Smad phosphorylation are commonly used to quantify BMP signaling (Humphreys et al., 2013).

All components of the BMP signaling module are members of multigene families that have evolved by duplication and divergence (Fritsch et al., 2010). In *Drosophila*, in addition to Dpp there are two other BMPs, Screw (Scw) and Glass bottom boat (Gbb), which are homologous to BMPs 5/6/7/8 in mammals. In addition to Thickveins (Tkv; ALK 3/6 in mammals), there are two other type I receptors: Saxophone (Sax; ALK 1/2 in mammals), and Baboon (Babo; ALK 4/5/7 in mammals). Conventional models of BMP signaling incorporate the one factor - one receptor - one function formula, with Dpp:Tkv:Punt:Mad comprising the canonical Dpp signaling module in the fly. Thus, while Tkv is thought to act primarily as a Dpp type I receptor, Sax is thought to transduce the Scw and Gbb signals. Punt, on the other hand, is thought to provide type II receptor function to complexes containing all Drosophila BMP family members (Dpp, Scw, and Gbb) (Gesualdi and Haerry, 2007). It is also well known that varying receptor components can affect signaling activity through interaction with different R-Smad family members (Haerry et al., 1998). However, while crosstalk between canonical (Dpp:Tkv:Punt:Mad) and non-canonical modules (e.g. Scw:Sax: Punt:Mad) is occasionally considered, with both the Scw and Gbb modules having been shown to play auxiliary roles in Dpp signaling by supplementing its activity (Arora et al., 1994a; Chen et al., 1998; Haerry et al., 1998; Khalsa et al., 1998; Nguyen et al., 1998), the potential versatility that can result from dual pathway activation by a single ligand has not been well studied.

Tight control of BMP signaling is vital for all processes, and the Dpp-dependent process of dorsal closure in the fruit fly is not an exception to this rule. Both loss- and gain-of-function Dpp signaling mutants disrupt dorsal closure, with each producing a distinctive phenotype. While loss of Dpp signaling activators leads to cuticular dorsal holes, loss of Dpp antagonists results in a gain-of-signaling phenotype characterized by hypotrophy of ventral denticle belts and puckering of the
dorsal midline (Bates et al., 2008). Mutants exhibiting the signature hyperactive Dpp signaling cuticular phenotype include raw, ribbon (rib), puckered (puc), and mummy (mmy) (Byars et al., 1999; Humphreys et al., 2013; Jack and Myette, 1997; Jack and Myette, 1999; Ring and Martinez Arias, 1993). raw, rib and puc all mediate their effects on Dpp secondarily via regulation of JNK signaling (Bates et al., 2008), but mmy’s effects on Dpp are direct (Humphreys et al., 2013).

Mmy, the pyrophosphorylase catalyzing the last step in the production of UDP-GlcNAc (Araujo et al., 2005), is required to attenuate epidermal Dpp signaling during dorsal closure (Humphreys et al., 2013) and points to a new inhibitory node for Dpp/BMP signaling - one that is dependent on GlcNAc. The mechanism by which Mmy regulates Dpp signaling is not immediately recognizable as GlcNAc contributes to a substantial fraction of the glycome. In this regard, post-translational modification by GlcNAc can occur by either N- or O-linkages and on nucleocytoplasmic, membrane-associated, or secreted proteins. GlcNAc is also integral to the synthesis of glycosylphosphatidylinositol, chondroitin sulfate, and heparin sulfate proteoglycans (Tonning et al., 2006).

Here, we show that the single Drosophila O-GlcNAc transferase (OGT) encoded by super sex combs (sxc) functions downstream of mmy to antagonize Dpp signaling in the Drosophila embryo. At the mechanistic level, we show that Dpp can signal via the BMP type II receptor Sax in vivo and that O-GlcNAcylation of this receptor regulates its activity. When O-GlcNAcylation is intact, Sax is silent and the BMP type II receptor Tkv transduces the epidermal Dpp/BMP signal. When O-GlcNAcylation is disrupted, Sax is activated and contributes to Dpp/BMP signaling in a manner that extends signaling range both temporally and spatially. While both Sax and Tkv require phosphorylation for activity, our data point to a previously unrecognized role for O-GlcNAc as a Sax (but not Tkv) inhibitor. Moreover, our data indicate that Sax is a direct target of Sxc, as Sax is post-translationally modified by O-GlcNAc in Sxc-dependent fashion. Finally, we show that Dpp/BMP signaling is a nutrient sensitive process, as maternal restriction of dietary sugar leads
to ectopic Dpp signaling and embryonic death. Thus, while BMP signaling regulators, acting on the pathway at all steps of signal transduction (extracellularly, at the membrane, in the cytoplasm, and in the nucleus) have been studied extensively (Yadin et al., 2016), to this list we can now add a cytoplasmic regulatory node that has the potential to serve as a nutrient-sensitive sensor for signaling.
RESULTS

The sxc-encoded OGT is required for embryonic development

mmy codes for the single UDP-N-acetylglucosamine pyrophosphorylase in Drosophila (Schimmelpfeng et al., 2006), and its requirement for attenuating epidermal BMP signaling during dorsal closure points to a previously unrecognized role for glycosylation in defining a restricted BMP activity field in the fly (Humphreys et al., 2013). UDP-GlcNAc serves as the precursor for a diverse set of glycosyl modifications that are catalyzed by a similarly diverse set of transferase enzymes (Lairson et al., 2008). With this knowledge as our foundation, we speculated that identification of a transferase(s) with a mmy-like loss-of-function cuticle phenotype would point us to the mechanism by which glycosylation regulates Dpp signaling.

Hundreds of enzyme glycosyltransferases (estimates range from 250 to 500) carry out protein glycosylation in vertebrates (Katoh and Tiemeyer, 2013; Schachter and Freeze, 2009), with more than 20 of these functioning downstream of Mmy to execute specific GlcNAcylation in the fly (Figure 1A). To identify the transferase(s) functioning downstream of Mmy in regulating Dpp/BMP signaling, we disrupted each of the predicted Drosophila β-1,3 glycosyl-transferases (Correia et al., 2003), as well as the single O-linked N-acetylglucosamine transferase (OGT) (Sinclair et al., 2009) by RNAi. To this end, we used the tubulin-Gal4 (tub-Gal4) driver to mediate ubiquitous expression of UAS-RNAi’s targeting each of the transferases. Analysis of cuticle phenotypes revealed that loss of the super sex combs-encoded OGT, via a UAS-RNAi transgene targeting sxc (Vienna stock #18611), results in a loss-of-function phenotype that is shared with mmy (Figure 1B-D).

Our discovery of sxc in a screen for embryonic lethality was somewhat unexpected given that sxc, despite a well-documented role as a regulator of Polycomb group proteins, has never been considered to be essential for embryogenesis in the fly (Gambetta and Muller, 2014; Ingham,
This said, \textit{sxc} is maternally deposited and expressed broadly throughout embryogenesis (Figure 2A-C), consistent with our RNAi studies pointing to a critical role for \textit{sxc} in development. Also, and although not published, anecdotal evidence for an early embryonic zygotic \textit{sxc} lethality has been noted (FlyBase, 2003).

In an effort to independently reproduce our RNAi findings, we conducted a lethal stage analysis of five \textit{sxc} mutants, documenting when and how \textit{sxc} mutants expire (Figure 1H). Two central findings emerged from this analysis: First, \textit{sxc} mutants suffer an incompletely penetrant lethality ranging from 20\% to 37\% for alleles in a series ranging from weakest to strongest (Figure 1I). Second, \textit{sxc} mutants exhibit an abnormal ventral cuticle analogous to that which we observed in \textit{sxctub}\textsuperscript{RNAi} studies, although we found that the extent of ventral denticle hypotrophy correlates with allele strength previously determined on the basis of lethality. In this regard, \textit{sxc}\textsuperscript{1} and \textit{sxc}\textsuperscript{6} mutant cuticles display no denticle belts (Figure 1E) while \textit{sxc}\textsuperscript{3}, \textit{sxc}\textsuperscript{5}, and \textit{sxc}\textsuperscript{7} cuticles display hypotrophied denticle belts (Figure 1F). We also showed that the cuticle phenotype of animals harboring the \textit{sxc}\textsuperscript{1} allele in \textit{trans} to a non-complementing deficiency is indistinguishable from that of \textit{sxc}\textsuperscript{1} homozygotes (Figure 1G), providing genetic evidence that the \textit{sxc}\textsuperscript{1} nonsense allele is, as previously suspected, null (Mariappa et al., 2018; Sinclair et al., 2009). Given that \textit{sxc}\textsuperscript{1} exhibits the greatest level of embryonic lethality, the strongest cuticle defect, and is genetically defined as null, we chose this allele for all further experiments. With respect to the discrepancy between our characterization of \textit{sxc} as an embryonic lethal and the longer history of characterization of \textit{sxc} as a pupal lethal, it is notable that \textit{sxc}-dependent embryonic lethality is an incompletely penetrant phenotype and that \textit{sxc}-dependent embryonic lethality can easily escape notice in preparations of devitellinized embryos.

Last, as we noted that \textit{sxc} is maternally deposited, we tested whether \textit{sxc} is required maternally for embryonic Dpp/BMP signaling. To do this, we used the maternal triple driver (Mazzalupo and...
Cooley, 2006) and RNAi to generate females lacking maternal sxc. Our examination of embryos deposited by females lacking maternal sxc showed no differences in cuticle appearance or viability in comparison to embryos derived from wild-type mothers. In contrast, embryos lacking both maternal and zygotic sxc contributions are indistinguishable from those lacking zygotic sxc only - both in terms of lethality and cuticle analysis (Figure 2D-L). Taken together, these data indicate that maternal sxc does not regulate embryonic Dpp/BMP signaling.

**Sxc antagonizes Dpp/BMP signaling**

Albeit incompletely penetrant, the dorsal-puckered, ventral-hypotrophied cuticle phenotype that we see in populations of sxc mutant embryos is a well-documented hallmark of ectopic Dpp signaling (Bates et al., 2008; Byars et al., 1999; Riesgo-Escovar and Hafen, 1997). Thus, we speculated that sxc-dependent O-linked glycosylation normally suppresses epidermal Dpp signaling in *Drosophila* embryos. We used two lines of experimentation to test this idea. First, we confirmed that the downstream transcription factor Mothers against Dpp (Mad), is required for the mispatterned cuticle that we observe in sxc mutants. In this regard, examination of cuticles deficient for both Mad and sxc revealed them to be identical to those derived from Mad alone (Figure 3A-B). Mad-dependent masking of the sxc phenotype points to a role for sxc as a Dpp/BMP signaling antagonist. Second, we directly compared epidermal Dpp activity in wild-type and sxc\(^1\) null embryos. In brief, we used an antibody directed against pMad (the phosphorylated [activated] form of Mad) in conjunction with confocal visualization methods. Using this system, we detected pMad very broadly in the epidermis of both wild-type and sxc embryos undergoing germ band extension. However, later in development, Dpp signaling, which normally wanes during dorsal closure in wild-type embryos, persists temporally and extends to greater depths in the epidermis of sxc embryos providing direct evidence that Sxc is a Dpp antagonist (Figure 3C-F).

Dpp signaling expansion in sxc mutants is identical to that which we documented previously in mmy\(^1\) mutants (Humphreys et al., 2013). In this regard, we observed that while pMad
immunoreactivity extends to a maximum average depth of five cells in the epidermis of wild-type embryos, immunoreactivity extends to a maximum average depth of ten cells in similarly staged sxc mutants (Figure 3G). The sxc pMad immunoreactivity phenotype, like the associated sxc lethality, is incompletely penetrant.

Antagonists of Dpp signaling during dorsal closure function as regulators of either the JNK or the Dpp/BMP signaling pathways. Three of the four antagonists we and others have studied thus far (raw, puckered [puc], and ribbon [rib]) function at the level of JNK (Byars et al., 1999). mmy, in contrast, targets the Dpp/BMP signaling pathway directly (Humphreys et al., 2013). None are part of the brinker (brk) repression system (Deignan et al., 2016; Minami et al., 1999). Results from three lines of experimentation are consistent with our hypothesis that sxc, like mmy, regulates Dpp/BMP signaling directly, rather than secondarily via JNK. First, Jun accumulates normally in LE cells in sxc mutants (Figure 3H-J, H'-J'). Second, Jun protein levels are not elevated in sxc mutants, particularly in comparison to a known JNK antagonist, raw (Figure 3K) (Humphreys et al., 2013). Third, the JNK reporter line, dpp^{151H} is properly expressed in the LE of sxc embryos (Figure 3L). It is also notable that the expression domain of the dpp transcriptional inhibitor brk is unaffected in sxc mutants (Figure 3M-N), as it is similarly unaffected in all Dpp/BMP signaling antagonists characterized to date (Humphreys et al., 2013).

Sxc antagonizes Dpp/BMP signaling via modulation of the Sax type I receptor

Having established that pMad persists broadly in the epidermis of sxc embryos despite normal signatures of JNK signaling, we next assessed requirements for Dpp pathway components in sxc-dependent epidermal Dpp signaling (Figure 4A). First, we tested whether ectopic Dpp signaling is dependent on LE dpp expression. To do this, we examined epidermal Dpp signaling in sxc Jra^{JA109} double mutants; Jra^{JA109} mutants do not express LE dpp, but leave all other embryonic dpp expression signatures intact (Humphreys et al., 2013). In comparison to sxc mutants where
epidermal Dpp signaling expands spatially and temporally (see Figure 3), we observed no Dpp signaling in sxc Jra^{A109} double mutants (Figure 4B-C). The absence of sxc-dependent Dpp signaling in the epidermis of sxc Jra^{A109} double mutants identifies LE dpp as the trigger for sxc-dependent Dpp signaling as it is also for wild-type Dpp signaling. In addition to its role in the epidermis, LE Dpp signaling induces heart formation in the embryonic mesoderm through activation of the tinman transcription factor (Lockwood and Bodmer, 2002; Xu et al., 1998). Ectopic Dpp signaling in sxc mutants is, however, restricted to the epidermis. We observed no embryonic heart (dorsal vessel) malformations, nor did we visualize any difference in tin expression in comparisons of sxc mutants with wild-types (Figure 4E-F).

Second, we considered the requirements for Punt and Tkv, the canonical receptors of the epidermal Dpp/BMP signaling ligand (Hamaratoglu et al., 2014) (see Figure 4A). Amorphic alleles of either punt (put) or tkv lead to a fully penetrant embryonic lethality due to defects in dorsal closure (Figure 4I, K). put^{68} and sxc^{1} put^{88} double mutants exhibit a shared loss-of-function dorsal-open phenotype demonstrating an essential epidermal role for put in both sxc-dependent and wild-type Dpp/BMP signaling (Figure 4J). Conversely, and somewhat to our surprise, tkv^{5} sxc^{1} double mutants exhibit a fully penetrant ectopic Dpp signaling cuticle phenotype (Figure 4L). Although quantitatively different (full rather than incomplete penetrance), the tkv^{5} sxc^{1} cuticle phenotype is qualitatively identical to that which we observed in sxc mutants (see Figure 1D-G), indicating that tkv in this context is not required for epidermal Dpp signaling. Given the unexpected nature of this finding, we confirmed that the tkv^{5} sxc^{1} cuticle phenotype is truly diagnostic of ectopic signaling by visualizing Dpp signaling directly. pMad stains are positive in tkv^{5} sxc^{1} double mutants as they are in sxc^{1} mutant embryos (Figure 4D). This ectopic signaling is not explained by differential expression of tkv, as tkv is similarly expressed in sxc^{1} and wild-type embryos (Figure 4G-H). Taken together, these data indicate that: 1) there is a type I receptor that transduces the Dpp/BMP signal via phosphorylation of Mad independently of Tkv, and 2) this type
I receptor activity is regulated by Sxc (OGT).

Last, we examined whether ectopic Dpp signaling observed in sxc mutants could be due to signaling through the other epidermal BMP type I receptor, Saxophone (Sax) (Figure 5I). We confirmed that sax is expressed ubiquitously in stage 15 embryos and that sax expression, like that of tkv, is not altered in sxc mutants (Figure 5A-C). Given that the spatio-temporal expression of sax positions it as a potential transducer of the epidermal Dpp signal, we tested whether ectopic Dpp signaling in sxc mutants requires sax. To do this, we analyzed Dpp signaling levels in sxc sax double, and sxc tkv sax triple mutants using amorphic alleles and assays of cuticle and pMad signaling. Dpp signaling levels in sxc\(^{1}\) sax\(^{5}\) embryos are comparable to wild-type in assays of cuticle phenotype and pMad signaling domains (Figure 5D-F), presumably because the Dpp signal is now funneled exclusively through Tkv. Consistent with this idea is our observation that all epidermal Dpp signaling is eliminated in the epidermis of triple mutants: tkv\(^{5}\) sxc\(^{1}\) sax\(^{5}\) embryos secrete a dorsal-open cuticle, indistinguishable from that of tkv\(^{5}\) or sax\(^{5}\)tkv\(^{5}\) double mutants and diagnostic of Dpp/BMP signaling loss (Figure 5G-H). We also assessed the function of Babo (the other Drosophila type I receptor) and Wishful thinking (Wit; the other Drosophila type II receptor) in double mutant studies. For this study, we disrupted babo using standard genetic means (the amorphic allele babo\(^{32}\)) and wit using RNAi gene disruption techniques (wit\(^{tub}\) RNAi). We found that both sxc babo and sxc wit double mutants secrete hypotrophic ventral cuticle (Figure 5J-K and data not shown). Thus, ectopic Dpp/BMP signaling requires neither babo nor wit, and the Sxc/OGT-dependent version of Dpp signaling is specific to the Punt:Sax receptor partnership. This result was not unexpected as Babo normally associates with dSmad2 (Smox in Drosophila) to transduce activins and Wit is expressed exclusively in the central nervous system and thought to transduce BMP signals there. This said, it is noteworthy that Babo exhibits a glucose repressive function that is independent of insulin in the Drosophila digestive tract (Chng et al., 2014).
Dpp signals via Sax as a homodimer.

Although Sax functions as a receptor for the Scw and Gbb members of the Drosophila BMP family (Twombly et al., 2009), our data suggest that Sax also functions as a receptor for Dpp (at least in the embryonic epidermis). Specifically, our discoveries that the sxc phenotype is: 1) dpp-dependent, but 2) only ~40% penetrant in a tkv\(^+\) background while fully penetrant in a tkv\(^{null}\) background suggests that Sax signals through the same ligand as Tkv (Dpp), albeit with an affinity for Dpp that is reduced in comparison to that of the Tkv receptor. Alternatively the Sax ligand might be a Dpp:Scw or Dpp:Gbb heterodimer, as BMP heterodimers have been demonstrated to have essential functions in vivo (Kunnapuu et al., 2014; Sawala et al., 2012; Shimmi et al., 2005).

We combined genetic and molecular methods to determine whether the bioactive Sax ligand is: 1) a Dpp homodimer, or 2) a heterodimer of Dpp in partnership with other BMPs, specifically Scw or Gbb (Raftery and Sutherland, 1999). First, we examined scw and gbb gene expression in whole mount embryos in situ. While scw and gbb are expressed ubiquitously in the early embryo, (Figure 6A-D), both genes are silenced in mid-embryogenesis (Figure 6E-H; FlyBase, 2003), and thus unlikely candidates for mediators of epidermal Dpp/BMP signaling. Consistent with their early expression, loss of either scw (in scw\(^{5}\) null heterozygotes) or gbb (in gbb\(^{D4}\) null heterozygotes) results in weak DV phenotypes (Figure 6 I, K) (Arora et al., 1994b; Harden, 2002). We generated sxc\(^{+}\) scw\(^{5}\) and sxc\(^{+}\) gbb\(^{D4}\) double mutants to assess whether disruption of either scw or gbb (like disruption of sax) restores wild-type Dpp signaling patterns to Sxc/OGT-deficient embryos. Cuticles derived from sxc\(^{+}\) scw\(^{5}\) and sxc\(^{+}\) gbb\(^{D4}\) double mutants exhibit hypotrophic denticle belts like those of sxc\(^{+}\) mutants (Figure 6J, L), indicating that neither Scw nor Gbb participates in Sxc-dependent Dpp/Put/Sax signaling. Taken together these genetic studies of ligands show that in the embryonic epidermis, Dpp signals through Sax as a homodimer and not as a heterodimer with either Gbb or Scw. Thus, despite the fact that Dpp can and does heterodimerize with other BMP
ligands (Anderson and Wharton, 2017; Hamaratoglu et al., 2014; Sawala et al., 2012), we found no evidence for such heterodimers in the embryonic epidermis during dorsal closure.

**Sax is modified by O-GlcNAc**

O-GlcNAcylation involves the transfer of a single β−GlcNAc moiety to serine (Ser) or threonine (Thr) residues of cytosolic or nuclear proteins, and in this regard O-GlcNAcylation is comparable to protein phosphorylation (Comer and Hart, 2000). Indeed, many OGT substrates are both O-GlcNAcylated and phosphorylated (potentially at several sites), sometimes even in reciprocal fashion. In other cases, O-GlcNAc modifies kinase accessibility to phosphorylation sites (Hart et al., 1995). Our genetic data suggest that Sax receptor activity is regulated by O-GlcNAc, and as Sax is a known target for phosphorylation by Ser/Thr kinases, we tested whether it is the direct target of Sxc in its modulation of epidermal Dpp/BMP signaling. To do this, we prepared lysates from wild-type and transgenic embryos (dorsal closure stage 8-12 hours after egg lay [AEL]) harboring a functionally-validated FLAG-tagged Sax isoform (Le et al., 2017). We extracted Sax-FLAG from transgenic embryo lysates by using ab13970 [AbCam], the antibody that recognizes FLAG. We assessed the O-GlcNAcylation state of the Sax-FLAG transgene in western blots of ab13970-affinity purified proteins by probing with RL2 (AbCam), an antibody that recognizes O-GlcNAc. O-GlcNAc-modification of Sax-Flag is dependent on Sxc, as glycosylation of Sax-FLAG does not occur in the presence of ubiquitously expressed RNAi targeting sxc (Figure 7A). In an extension of this analysis to a functional GFP-tagged version of Tkv (Hsiung et al., 2005), we showed that the Tkv type I receptor is not modified by O-GlcNAcylation (Figure 7A), consistent with our genetic prediction that Sxc only inactivates Sax.

**Dpp signaling is sensitive to glucose availability**

Throughout the life histories of metazoans, longevity and fertility are heightened in favorable
energetic conditions (e.g. food abundance) and diminished in unfavorable conditions (e.g. food scarcity) (Fontana and Della Torre, 2016; Panth et al., 2018). In accordance with these associations, our observation that O-GlcNAc normally suppresses Sax-mediated transduction of Dpp evokes the expectation that nutrient-poor conditions will activate Sax-mediated transduction of Dpp/BMP - presumably to augment Dpp/BMP signaling and limit embryo viability in sub-optimal growth conditions. To determine whether *Drosophila* manifest diet-mediated linked effects on fertility and Dpp signaling, we assessed viability and Dpp signaling in embryos derived from mothers fed varying amounts of glucose. We used sucrose as the dietary source of glucose: (no sucrose diet (NSD), low sucrose diet (LSD; 0.15 M sucrose), standard sucrose diet (SSD; 0.5 M sucrose), and high sucrose diet (HSD; 1.0 M sucrose) (Chng et al., 2014; Piper et al., 2014). In this dietary study, we observed lethality measures in NSD conditions comparable in quantity (40%) to those observed in *sxc* embryos (Figure 7B). In addition, cuticles of embryos derived from wild-type females raised on no-sucrose diets exhibit a shared loss-of-function phenotype with *sxc* mutants, suggestive of ectopic Dpp signaling in the embryonic epidermis (Figure 7C-D). In an extension of this analysis to *sxc* mutants, we showed that depletion of maternal dietary sugar does not enhance lethality in *sxc* mutant embryos (Figure 7B) or affect phenotype (Figure 7E). Moreover, depletion of sugar from the diet of *sax* mothers fully rescues viability and restores normal cuticle phenotype (Figure 7B, F). Taken together, these data demonstrate that embryonic Sax-mediated Dpp signaling is sensitive to maternal sugar intake, and that Sxc is the sugar sensor.
DISCUSSION

Although several well-characterized cellular regulatory systems are known to fine-tune the amplitude and range of Dpp/BMP signaling (Bollenbach et al., 2008; Humphreys et al., 2013; Raftery and Umulis, 2012; Schwank et al., 2011), how the pathway responds to environmental cues has remained largely unexplored. Here we describe a previously unrecognized cytoplasmic Dpp/BMP signaling regulatory node that relays environmental nutritive conditions to the developing embryo. Building on our previous demonstration that UDP-GlcNAc is a regulator of Dpp signaling in the Drosophila epidermis (Humphreys et al., 2013), we have now defined the mechanism by which UDP-GlcNAc regulates this pathway. It is through Dpp type I receptor O-GlcNAcylation. When O-GlcNAcylation signals are intact (sxc+), Dpp signals only in the most dorsal regions of the epidermis. In contrast, when O-GlcNAcylation marks are disrupted (sxc−), the Dpp signaling range is extended with respect to both space and time (see Figures 1 and 3).

As expected for a modulator of Dpp signaling, Sxc mediates its effects via Dpp, its canonical type II receptor Punt and its R-Smad intracellular transducer Mad (see Figure 4). However, in an unexpected but revealing twist, we found that Sxc modulates Dpp signaling not via its canonical type I receptor Tkv, but rather via the homologous type I receptor Sax (see Figure 5). Moreover, we found that Sax is O-GlcNAcylated by Sxc (see Figure 7). Taken together, our studies show that both Tkv and Sax function as Dpp receptors, extending previous views of Sax as a receptor dedicated to the other BMP-type ligands in Drosophila (Scw and Gbb) (Nguyen et al., 1998). Our studies also show that despite their association with the same R-Smad (Haerry, 2010), the Tkv and Sax responses to Dpp are substantively different, with spatial and temporal output properties distinguishing the two. Finally, our studies indicate that sxc is the genetic toggle that modulates Dpp:Sax pathway output and that Sax is the O-GlcNAc modified target of Sxc.
The Sax family of Type I BMP receptors

The fundamental processes of gene duplication and divergence generate protein families with functionally versatile members. The BMP type I receptor family is an example of one family that has evolved by duplication and divergence. In *Drosophila*, there are three BMP type I receptors: Tkv (ALK 3/6 in mammals), Sax (ALK 1/2 in mammals), and Babo (ALK 4/5/7 in mammals). Tkv is the best characterized of the *Drosophila* type I ALK receptors, having several very well documented roles in *Drosophila* development. Tkv is thought to act primarily as a Dpp receptor (BMP 2/4 in vertebrates), while Sax, first identified in the Heidelberg screens for mutants disrupting cuticle pattern (Schupbach and Wieschaus, 1989), has been thought to function solely in response to the other two BMP-related ligands in Drosophila: Screw (Scw) and Glass bottom boat (Gbb), representing BMPs 5/6/7/8 in mammals (Haerry et al., 1998). In some developmental contexts, multiple BMP ligands are expressed in the same regions and thus require multiple matching receptors to transduce signals from all BMP ligands present. In the early embryo, for example, Scw, Gbb, and Dpp are expressed in the dorsal half of the embryo. While Dpp is the major determinant of dorsal fates, Scw and Gbb have been shown to play auxiliary roles in BMP signaling by supplementing Dpp activity. (Arora et al., 1994a; Chen et al., 1998; Haerry et al., 1998; Khalsa et al., 1998). In this respect, maternal loss of the ligands Scw/Gbb or their type I receptor Sax results in a mild ventralizing phenotype, whereas maternal loss of Dpp or its type I receptor Tkv results in a completely ventralized embryo (Arora et al., 1994b; Irish and Gelbart, 1987; Neul and Ferguson, 1998; Twombly et al., 2009). In dorsoventral axis determination Sax transduces the Scw signal, thereby augmenting but not directly facilitating Dpp signaling via Tkv (Nguyen et al., 1998).

Years of additional molecular and genetic studies helped shape our mechanistic understanding of Sax as a type I BMP receptor (Anderson and Wharton, 2017; Le et al., 2018; Wharton and Derynck, 2009). In the wing imaginal disc, Sax functions to transduce Gbb/BMP signaling directly
in cooperation with Tkv (Haerry et al., 1998); and can also limit the availability of the Gbb ligand, thus acting as a Gbb/BMP signaling antagonist (Bangi and Wharton, 2006). Both functions of Sax help control wing growth and patterning via transduction of Gbb signals, suggesting that precise control of Sax activity is critical. While we have long understood that Sax transduces the Scw and Gbb signals, in this study, we have identified a function for Sax in transducing the Dpp signal in the epidermis independently of Tkv (see Figure 5).

Tkv is the most prominent of the Drosophila BMP type I receptors, with several well-defined developmental roles. Tkv is part of the canonical Dpp signaling module (Dpp:Tkv:Punt:Mad) that is required for Drosophila embryonic viability. The pathway is required maternally for dorsal axis formation and zygotically for dorsal closure. Given the breadth and depth of genetic sax studies that precede this one, our discovery that Sax can, like Tkv, transduce an epidermal Dpp signal was somewhat unexpected. However, we have also found that epidermal Sax activity is normally suppressed by O-GlcNAcylation (see Figure 7), and this observation is consistent with genetic loss-of-function studies revealing normal dorsal closure and embryonic viability in animals homozygous for null alleles of sax (Twombly et al., 2009). Importantly, our demonstration that Dpp signals via Sax extends conventional signaling models, which incorporate the one factor - one receptor - one function formula, with Dpp:Tkv:Punt:Mad comprising the canonical Dpp signaling module in Drosophila. In this regard, we show that Dpp can also signal in a Dpp:Sax:Punt:Mad module, but that this module is inactive (or the receptor complex does not form) in conditions of readily available glucose (i.e. normal diet). While crosstalk between canonical and non-canonical modules (e.g. Scw:Sax:Punt:Mad) has occasionally been considered in vitro (e.g. (Haerry et al., 1998), the potential versatility that can result from dual pathway activation by a single ligand has not been well studied.

Finally, our discovery that Sax activity in sxc mutants is: 1) dpp-dependent, but 2) only ~40%
penetrant in a tkv\textsuperscript{*} background while fully penetrant in a tkv\textsuperscript{null} background indicates that Sax and Tkv signal through the same ligand (Dpp) in vivo, albeit with inherently different properties. This finding is consistent with functional studies of the two receptors in culture, where measures of Tkv activity are 10 fold greater than those of Sax (Haerry, 2010). This said, at least three receptor properties, none mutually exclusive, might contribute to differential type I receptor signaling capacities: 1) ligand/receptor interactions; 2) receptor kinase enzyme kinetics and 3) receptor expression domains. Indeed, differences in each of these properties are natural consequences of gene duplication and evolution. Further probing of receptor subtype-specific variations will provide insight into key selectivity determinants for individual responses to ligand (Haerry, 2010; von Bubnoff and Cho, 2001).

**The Sax family of Type I Dpp/BMP receptors in human health and disease**

As we have found in to be the case for Drosophila Sax, disease and developmental phenotypes in humans are associated with activating mutations of the human Sax orthologue Activin A Receptor (also called Type II-Like 1 and 2, ACVRLK1/2, and ALK1/2). Mutations of human Activin A Receptor, are causative of Fibrodysplasia Ossificans Progressiva (FOP; OMIM ID#135100; de la Pena et al., 2005). FOP is an autosomal dominant disorder characterized by widespread ossification of soft tissue and occurs sporadically throughout life, with most patients wheelchair bound by their third decade (Petrie et al., 2009). The R206H amino acid substitution, which is responsible for most FOP occurrences (Le and Wharton, 2012), is activating in cell culture, a result of ligand-independent and constitutive ACVR1 phosphorylation (de la Pena et al., 2005). Disease resulting from ectopic activation of ACVR1 argues that precise control of Activin A Receptor activity (like that of Sax) is critical for proper signaling during adulthood, and aligns with data herein demonstrating that precise control of Sax receptor activity is also important for embryonic development.
An embryonic role for Sxc and O-GlcNAcylation

There is a single *Drosophila* OGT-encoding gene, *sxc*, and it was first identified based on its loss-of-function phenotype - ectopic sex combs on the second and sometimes third leg pairs of pharate adult males (Ingham, 1984; Sinclair et al., 2009). Here, we demonstrate that *sxc* mutants additionally suffer an incompletely penetrant embryonic lethality associated with defects in Dpp signaling (see Figure 1). Developmental roles for *sxc* homologs have also been described in vertebrates, where the gene exists in single copy as well. In mice, amorphic *Ogt* mutations are associated with embryonic lethality (O'Donnell et al., 2004). Mutations of the human *OGT*, which are presumed to be hypomorphic, are embryonic viable but lead to X-linked intellectual disability (Willems et al., 2017; Pravata et al., 2019)

OGT modifies cytosolic and nuclear proteins (Bond and Hanover, 2015). In contrast to N-glycosylation, which occurs in the Golgi or ER and produces a variety of sugar modifications in chains and branches, O-GlcNAcylation occurs in the cytoplasm as a reversible addition of a single GlcNAc residue onto serine or threonine residues. In this regard O-GlcNAcylation is comparable to protein phosphorylation (Hart et al., 2011; Hu et al., 2010). Indeed, proteins can be both O-GlcNAcylated and phosphorylated, even at several sites. However, at any given site, modifications by O-GlcNAc and phosphorylation are mutually exclusive (Leney et al., 2017). This switch provides a mechanism by which O-GlcNAcylation can inactivate a protein normally activated by phosphorylation. Notably, our data show that Sax, but not Tkv, is O-GlcNAcylated by Sxc.

Bioinformatic survey of BMP type I receptors, using the YinOYang 1.2 algorithm (Gupta and Brunak, 2002) reveals pairs of high probability OGT modification sites in Sax/ALK1/2 type I receptor family members (e.g. T228 and S229 in SaxPA and S159 S160 in ALK1) that are not conserved in Tkv/ALK 3 family members, in addition to three sites in the GS domain that are
conserved in both Sax and Tkv (T244, S245, and T259 in SaxPA) (Figure 7I) (Gupta and Brunak, 2002). All high probability targets map within or immediately adjacent to the GS domain where Ser/Thr phosphorylation is required for type I receptor kinase activation (and where higher levels of phosphorylation correspond to greater kinase activity (Wieser et al., 1995). Moreover, this is the domain that harbors receptor activating mutations of FOP (R206H in ACVR1; Shore et al., 2006). Our study, along with a recent report of O-glycosylation-dependent inactivation of the Gbb BMP-type ligand (Anderson and Wharton, 2017), points to post-translational modification by O-glycans as a critical component of cell signal regulation in development and disease (Anderson and Wharton, 2017).

As the identification of diverse sets of O-GlcNAcylated proteins advances, it becomes increasingly clear that O-GlcNAc is an abundant modification with important implications not only for cell signaling, but also for protein function more generally. Important signaling molecules such as Protein kinase C, extracellular signal-regulated kinase, Runx2, CCAAT/enhancer-binding proteins have all been shown to be modified by O-GlcNAcylation (Sun et al., 2016) and we can now add Sax to this ever-expanding list. Finally, there are notable similarities between our studies of Dpp signaling in whole animals and a recent study of Hippo signaling in a cultured cells. In the case of the latter, the Hippo signal transducer, YAP was shown to be O-GlcNAcylated. Moreover, this modification prevented YAP phosphorylation and thus repressed its transcriptional activation function (Peng et al., 2017). The authors of this study also demonstrated that the process is regulated by glucose in the culture medium, thereby linking environmental nutrition to signal activation and transcriptional changes.

**Nutritional sugar**

Given the potential for complex interactions between O-GlcNAcylation and phosphorylation (Comer and Hart, 2000), glucose offers a potential target for controlling development in poor
nutrient conditions and/or tissue homoeostasis and regeneration in ageing and disease. Emerging
evidence suggests that altered levels of glucose influence osteogenic, chondrogenic and
adipogenic differentiation via the insulin, TGF-β and peroxisome proliferator-activated receptor
pathways, among others (Sun et al., 2016). Our data provide insight into the mechanism by which
OGT loss impacts signaling by Dpp, a member of the TGF-β superfamily of secreted cytokines.
Albeit incompletely penetrant, a high proportion of animals harboring amorphic alleles of sxc (the
Drosophila gene encoding OGT) suffer an embryonic lethality due to ectopic epidermal Dpp
signaling. Notably, this phenotype is mimicked in a similar fraction of embryos derived from
females deprived of dietary sugar. Thus, deficiencies in the hexosamine biosynthetic pathway,
and a lack of O-GlcNAc production in particular, mimics the loss of sxc in the fly embryo.
Importantly, this phenotype is not observed in sax embryos demonstrating that the lethality and
Dpp phenotypes induced by the lack of dietary sugar requires Sax activity (see Figure 7).

Extracellular glucose flux modulates intracellular O-GlcNAc levels through the hexosamine
biosynthetic pathway, with increased glucose uptake leading to increased production of cellular
GlcNAc, the OGT substrate (Zachara and Hart, 2004a, b). Several lines of evidence, most notably
in studies of patients afflicted with diabetes, have led to the speculation that OGT functions as a
nutrient sensor (Konrad and Kudlow, 2002; Majumdar et al., 2004; Schwartz and Pirrotta, 2009;
Bond and Hanover, 2015). In this regard, a role for OGT as a nutrient sensor could have major
impacts on multiple cellular processes, including signaling by Dpp, a member of the TGF-β
superfamily of secreted cytokines with multiple developmental roles. Our data demonstrate that
complete loss of sxc, which codes for the single OGT in Drosophila, results in an embryonic
lethality associated with ectopic Dpp signal activation in the epidermis. This phenotype is
mimicked in the offspring of mothers deprived of dietary sugar, suggesting that a lack of HBP
input (low sugar) and the resultant lack of HBP output (low O-GlcNAc) is equivalent to genetic
loss of sxc in fly embryos. Importantly, gain-of-function Dpp phenotypes that are induced by a
lack of dietary sugar in wild-type embryos are not observed in sax embryos, thus demonstrating that Sax activity modulates this dietary response (Figure 7).

In conclusion, our data point to a Sax-mediated branch of epidermal Dpp signaling that is normally inhibited by the hexosamine biosynthetic pathway product O-GlcNAc. Moreover, our data bolster recent suggestions that the hexosamine biosynthetic pathway might mediate crosstalk between glucose flux, cell signaling, cell differentiation, and organismal development.

ACKNOWLEDGEMENTS

The authors thank Sandra Kazuko for technical support, John Hanover, Tom Kornberg, and Kristie Warton for fly lines, and Diana Lim for figure preparation.

AUTHOR CONTRIBUTIONS


M.J.M. and A.L. conceived and designed the experiments, analyzed the data, and wrote the paper. A.L. and G.B.H. conceived of the RNAi screen and G.B.H. carried out the screen. M.J.M. performed all other experiments, with help from A.K. in Western analyses. A.L. supervised the project.

DECLARATION OF INTEREST

The authors declare no competing interests.
Figure Titles and Legends

Figure 1. sxc mutations result in embryonic lethality
(A) UDP-GlcNAc synthetic pathway and downstream utilization of GlcNAc. (B-G) Images of embryonic cuticle with genotypes indicated. (H) sxc mutants; the sxc$^3$ lesion does not map to the sxc locus and the molecular nature of the lesion has not been defined. (I) Quantification of embryonic lethality in different sxc mutants (**p<0.01, n=>95/genotype, mean ± SD). In this and all subsequent figures, embryos are viewed laterally, with anterior to the left.

Figure 2. Zygotically-derived sxc modulates development
(A-I) Hybridizations in situ to visualize embryonic sxc expression. Row and column headers indicate embryonic genotypes and stages. (J-K) Images of embryonic cuticle with genotypes indicated. (L) Quantitation of embryonic lethality associated with maternal, zygotic, and maternal/zygotic loss of sxc (**p<0.01, n=>200/genotype, mean ± SD). Maternal is abbreviated Mat, and Tubulin is abbreviated Tub.

Figure 3. Aberrant Dpp signaling in sxc mutants is not due to JNK signaling defects
(A-B) Images of embryonic cuticle with genotypes indicated. (C) Model of JNK-induced Dpp signaling in wt and sxc$^1$ embryos. In stage 12-13 embryos JNK-dependent dpp expression in leading edge cells (half-filled circles) triggers Dpp signaling broadly in the lateral epidermis (to an average depth of five cells in the epidermis of wild-type embryos and to an average depth of ten cells in sxc mutants; filled circles). Dpp signaling wanes in stage 14-15 wild-type embryos but persists in sxc$^1$ embryos. (D-F) pMad stains of stage 14 embryos with genotypes indicated. (G) Quantification of depth of pMad staining nuclei in wild-type and sxc$^1$ embryos (n=>10/genotype, mean ± SD). (H-J) Jun protein localization in embryos with genotypes indicated. (H'-J') Enlargement of boxed areas in H-J. (K) Western blot analysis of Jun protein in lysates from wild-
type and mutant embryos (stage 12-15). JNK reporter expression in sxc\(^{1}\) mutant embryos, stage 12-15). (M-N) Hybridizations \textit{in situ} to visualize embryonic expression of the Dpp antagonist; genotypes are indicated.

**Figure 4. sxc-dependent Dpp signal transduction does not require Tkv**

(A) Schematic of the canonical Dpp receptor complex (Put and Tkv), with signal transduced intracellularly through the action of pMad. The constitutively active Put serine/threonine kinase activates Tkv via phosphorylation. Upon phosphorylation, the Tkv serine/threonine kinase activates Mad. (B-D) pMad stains of stage 12-13 embryos with genotypes indicated. (E-H) Hybridizations \textit{in situ} to visualize \textit{tin} and \textit{tkv} expression in stage 12-13 wt and sxc\(^{1}\) mutant embryos. (I-L) Images of embryonic cuticle with genotypes indicated.

**Figure 5. sxc-dependent Dpp signal transduction requires Sax**

(A-C) Hybridizations \textit{in situ} to visualize embryonic sax expression in stage 12-13 embryos with genotypes indicated. \textit{sax} is expressed in the epidermis of wt embryos, and its expression is not altered in similarly staged sxc mutants. Sax detection is specific as it is absent in control embryos expressing RNAi targeting \textit{sax}. (D-E, G-H, J-K) Images of embryonic cuticle with genotypes indicated. (F) pMad staining in sxc\(^{1}\) sax\(^{5}\) double mutants. (I) Schematic of the Dpp/Put/Sax receptor complex module that is active in sxc mutants, with signal transduced intracellularly through the action of pMad. The constitutively active Put serine/threonine kinase activates Sax via phosphorylation. Upon phosphorylation, the Sax serine/threonine kinase activates Mad.

**Figure 6. sxc-dependent Dpp signal transduction does not require scw or gbb**

(A-H) Hybridizations \textit{in situ} to visualize gene expression in embryos with genotypes identified in the lower left of each panel. Row and column headers identify hybridization probes (either scw
or gbb) and embryonic stages (either 5 or 12-13). (I-L) Images of embryonic cuticle with genotypes indicated.

**Figure 7. Dpp signaling through Sax is modulated by O-linked glycosylation and dietary sugar**

(A) Western blot analysis of O-GlcNAcylation status of Sax and Tkv. (B) Embryonic lethality quantification of embryos from adults raised on varying amounts of dietary sugar (**p<0.01, n=>100/genotype, mean ± SD). (C-F) Images of embryonic cuticle with genotypes indicated. (G) Schematic showing effect of Sxc-dependent glycosylation of Sax on Dpp receptor complex function. Although glycosylation clearly blocks receptor complex activity; our data do not allow us to determine whether complex assembly is inhibited by glycosylation. Yellow highlights mark predicted O-GlcNAcylation sites in Sax and Tkv and in their human homologs.
STAR METHODS

CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Anthea Letsou (aletsou@genetics.utah.edu).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Animals

Drosophila melanogaster were raised on a standard cornmeal diet. Embryos were collected using standard fly laying blocks and agar/grape juice plates supplemented with yeast. In the case of dietary sugar alteration, grape juice was omitted from the agar plates and sucrose (at concentrations indicated) was added to the standard yeast paste diet.

METHOD DETAILS

Fly strains

The Oregon R strain served as the wild-type in all experiments. Stocks were obtained from the Bloomington Drosophila Stock Center and the Vienna Drosophila Resource Center (see Key Resources Table). The sxc\textsuperscript{2}, sxc\textsuperscript{3}, sxc\textsuperscript{4}, sxc\textsuperscript{5}, sxc\textsuperscript{7} mutants were a gift of John Hanover (NIH). Double mutant analysis was performed using the sxc\textsuperscript{7} null allele and null alleles of all other genes unless otherwise noted in the text. The dpp\textsuperscript{151H}, tkv\textsuperscript{5}, and put\textsuperscript{88} mutants have been described (Johnson et al., 2003; Terracol and Lengyel, 1994) (Simin et al., 1998). UAS:Sax-Flag and UAS:Tkv-GFP lines were the generous gift of Kristi Wharton (Brown University) and Tom Kornberg (UCSF), respectively.

Phenotypic analysis

Cuticles were prepared using standard methods (Humphreys et al., 2013). In brief, embryos were dechorionated in 50% bleach and devitellinized in a 1:1 methanol/heptane mixture and afterwards
incubated in 1-step mounting media (100 ml Glacial Acetic Acid; 50 ml CMCP10; 25 ml 85% Lactic Acid) overnight at 65 degrees C. Cuticles were visualized via dark field microscopy on a Zeiss Axioskop. Lethal stage analyses were performed by plating embryos on grape juice agar plates and monitoring viability 48 AEL (after egg lay).

Immunohistochemistry and RNA in situ hybridization staining procedures were performed using standard methods (Humphreys et al., 2013). In brief, rabbit anti-phospho-Smad1,5 Ser463/465 (1:20, Cell Signaling Technology) or digoxigenin-labeled anti-sense RNA probes were incubated overnight on fixed embryos. Embryos were incubated overnight with secondary antibodies targeting pMad (goat anti-rabbit alkaline phosphatase [Jackson ImmunoResearch] or goat anti-rabbit Alexa Fluor 488 [Invitrogen Molecular Probes]), or digoxigenin-labeled RNA probes (mouse anti-digoxigenin alkaline phosphatase Fab fragments [Roche]). The following day, alkaline phosphatase detection was performed using nitro blue tetrazolium chloride and 5-Bromo-4-chloro-3-indolyl phosphate followed by brief dehydration in methanol and overnight incubation in 80% glycerol. All images were captured using a Zeiss Axioskop microscope with DIC optics.

**Protein studies**

Western blot studies were performed using protein lysates isolated from embryos 8-12 hrs AEL. Lysate samples were loaded onto a 12% Acryl-Bis polyacrylamide gel and subjected to electrophoresis at 100v for 3 hours. The gel was transferred to a PVDF membrane (Millipore) and blocked using 5% milk or 5% BSA in TBS + 0.05% Tween for 2 hours. Primary antibodies were used at 1:200 (anti-Jun) or 1:1,000 concentrations (anti-O-GlcNAc [RL2] from AbCam and anti-Flag [M2] from Sigma-Aldrich). HRP-conjugated goat-anti-mouse IgG secondary was used at 1:100,000 for anti-O-GlcNAc and 1:15,000 for anti-Flag, as well as HRP-conjugated goat anti-rabbit secondary at 1:5000 for anti-Jun. Blot detection was performed using equal volumes of
ECL Luminol solutions A and B (Santa Cruz) and imaged on a Mini-Medical Series machine (AFP Imaging).

Immunoprecipitation assays were performed on non-denatured lysates isolated from embryos 8-12 hrs AEL. 3 μg of antibody (anti-Flag or anti-GFP) was incubated with lysates for 90 minutes followed by a 60-minute incubation with protein-G sepharose beads (Santa Cruz). Beads were pelleted by centrifugation at 8,000 x G for 7 minutes. Supernatant was collected, and pelleted beads were subsequently washed in 500 μl lysis buffer 3 times. 2X Laemelli Sample Buffer was added to pellet and supernatant fractions, which were used subsequently in Western blot studies.

QUANTIFICATION AND STATISTICAL ANALYSIS

pMad Quantification

Quantification of pMad staining was performed by counting pMad positive nuclei of segments T1, T3, A4, and A6 within single image slices. The average number of cells from all quantified segments was used as the value for each animal tested (wt n=6, sxc<sup>1</sup> n=14).

Statistical Analysis

Lethal phase and pMad quantification datasets were assembled in Microsoft Excel 2013 and pairwise t-tests were performed with a statistical significance cutoff at p<0.05.
REFERENCES


Figure 2
Moulton & Letsou
Figure 7
Moulton & Letsou
**KEY RESOURCES TABLE**

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### Critical Commercial Assays

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### Experimental Models: Organisms/Strains

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<td>D. melanogaster: Tubulin Gal4 driver: y&lt;sup&gt;1&lt;/sup&gt; w*; P{tubP-GAL4}LL7/TM3, Sb&lt;sup&gt;1&lt;/sup&gt; Ser&lt;sup&gt;1&lt;/sup&gt;</td>
<td>Bloomington Drosophila Stock Center</td>
<td>RRID:BDSC_5138</td>
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<td>D. melanogaster: RNAi line of sxc</td>
<td>Vienna Drosophila Resource Center</td>
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<td><strong>D. melanogaster:</strong> RNAi line of sxc</td>
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<td><strong>D. melanogaster:</strong> allele of dpp: EcoI\lacZ&lt;sup&gt;dpp-151H&lt;/sup&gt;</td>
<td>Johnson et al., 2007, Genetics</td>
<td>FlyBase: FBal034426</td>
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<td><strong>D. melanogaster:</strong> allele of tkv: tkv&lt;sup&gt;5&lt;/sup&gt;/CyO</td>
<td>Bangi, E. and Wharton, K., 2006, Development</td>
<td>FlyBase: FBal0016822</td>
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<td><strong>D. melanogaster:</strong> allele of put: put&lt;sup&gt;68&lt;/sup&gt;/TM3</td>
<td>Simin, et al., 1998, Genetics</td>
<td>FlyBase: FBal0044294</td>
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<td><strong>D. melanogaster:</strong> sax-Flag expressing line: sax[Scer\UAS.T:Zzzz\FLAG]</td>
<td>Le, V.Q. and Wharton, K., 2012, Developmental Dynamics</td>
<td>FlyBase: FBal0268552</td>
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<td><strong>D. melanogaster:</strong> tkv-GFP expressing line: Scer\UAS.tkv-GFP</td>
<td>Hsuing, et al., 2005, Nature Letters</td>
<td>FlyBase: FBal0177447</td>
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**Oligonucleotides**

| **Recombinant DNA**

| **Software and Algorithms**

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