Molecular mechanism and functional significance of acid generation in the *Drosophila* midgut

Gayle Overend, Yuan Luo, Louise Henderson, Angela E. Douglas, Shireen A. Davies & Julian A. T. Dow

The gut of *Drosophila melanogaster* includes a proximal acidic region (~pH 2), however the genome lacks the H⁺/K⁺ ATPase characteristic of the mammalian gastric parietal cell, and the molecular mechanisms of acid generation are poorly understood. Here, we show that maintenance of the low pH of the acidic region is dependent on H⁺ V-ATPase, together with carbonic anhydrase and five further transporters or channels that mediate K⁺, Cl⁻ and HCO₃⁻ transport. Abrogation of the low pH did not influence larval survival under standard laboratory conditions, but was deleterious for insects subjected to high Na⁺ or K⁺ load. Insects with elevated pH in the acidic region displayed increased susceptibility to *Pseudomonas* pathogens and increased abundance of key members of the gut microbiota (*Acetobacter* and *Lactobacillus*), suggesting that the acidic region has bacteriostatic or bacteriocidal activity. Conversely, the pH of the acidic region was significantly reduced in germ-free *Drosophila*, indicative of a role of the gut bacteria in shaping the pH conditions of the gut. These results demonstrate that the acidic gut region protects the insect and gut microbiome from pathological disruption, and shed light on the mechanisms by which low pH can be maintained in the absence of H⁺, K⁺ ATPase.

Many animals generate a gut region of low pH to aid protein digestion, absorption of nutrients such as calcium, iron and vitamin B12, and to kill orally-acquired gut pathogens and parasites. The mechanism of acid generation in the human stomach is understood in considerable detail. Hydrochloric acid is generated in parietal cells via a proton pump (H⁺, K⁺-ATPase, a member of the P-type ATPase family, which expends ATP to generate an ion gradient across a membrane). The release of gastric acid is modulated by paracrine, endocrine or neural receptors, and is stimulated by food consumption. H⁺, K⁺-ATPase function is essential for normal secretion of gastric acid, and it has become a target for drugs such as Omeprazole, which treat excess acidity by binding non-competitively to disrupt acid generation. In addition to the H⁺, K⁺-ATPase (ATP4A); potassium channels, chloride transporters, and members of the Na⁺, K⁺-ATPase and NHE families are all vital for ion movement. A carbonic anhydrase is also essential for the rapid generation of protons and bicarbonate, which is exchanged for chloride by a Cl⁻/HCO₃⁻ anion exchanger on the basal membrane. Together, these transporters recycle Na⁺ and K⁺ across the basal and apical membranes of the parietal cell, resulting in the generation of gastric acid through release of H⁺ and Cl⁻ into the gastric gland lumen.

Intriguingly, the gut of *Drosophila melanogaster* includes an anterior acidic region (~pH 2), analogous to the vertebrate stomach; but there is no evidence for a homologue of the H⁺, K⁺-ATPase in the *Drosophila* genome, or indeed beyond the Boreoeutherian mammals (NCBI Homologene: http://www.ncbi.nlm.nih.gov/homologene/68081). This suggests that insects have evolved a novel mechanism for gastric acid pH generation that circumvents the requirement for direct exchange of protons for potassium ions; as insects make up the majority of living species on Earth, this is a significant issue. The acidic region of *Drosophila* midgut contains a unique subset of cells—the copper cells—with a highly invaginated apical membrane, similar to the mammalian gastric parietal cells. Each copper cell is bordered by a septate junction with an interstitial cell, which may also be involved in...
in acid pH generation. In the developing larva, the homeotic gene labial is essential for the specification of functional copper cells\(^\text{12,13}\); however, the ion transporters involved in pH generation have not been characterized, and the significance of the acid region in shaping interactions with the gut microorganisms\(^\text{14–16}\) has not been investigated. In adult however, recent studies have linked aging to epithelial metaplasia of the midgut, which can result in loss of gut pH and changes to the composition and load of intestinal microbiota\(^\text{17}\), suggesting a continuing role for gut pH throughout the lifespan of the fly.

The aim of this study was to characterize the mechanism of acid production in \textit{Drosophila} midgut, and to investigate its functional significance. We show that acid pH generation in the \textit{Drosophila} midgut requires input from at least six proteins or complexes with roles in ion homeostasis. We demonstrate that the larval acidic region plays a vital role in the ion homeostasis of the animal, confers protection against a bacterial pathogen, and controls the populations of gut bacteria.

### Results and Discussion

**The \textit{Drosophila} larval midgut is defined by five regions of pH.** Previous studies reported that the larval midgut contains at least four defined regions of different pH along its length\(^\text{10}\). By maintaining larvae on diet that contains a range of dyes with distinct pH transition points, a map of the pH in each gut region was obtained (Fig. 1). Five discrete regions of pH were resolved; the anterior midgut (pH 7), the acidic region (pH 2), the neutral region (pH 7), the transitional region (pH 6) and finally the posterior alkaline region (pH 9.5) (Fig. 1B). These five segments of the larval gut map approximately to the major segments defined in the adult\(^\text{18}\). Although the anterior and acidic regions are spatially distant from the posterior alkaline region in the excised midgut (Fig. 1B), they are closely apposed in the intact larva (Fig. 1A).

**Expression of the H\(^+\) V-ATPase complex in anterior midgut is essential for acid pH generation.** Given that the \textit{Drosophila} genome lacks an annotated H\(^+\), K\(^+\) ATPase, we sought to identify other transporters, for example the V-ATPase, that might play a key role in acidification\(^\text{13,19}\). Accordingly, we interrogated our RNAseq regional expression atlas for the larval midgut (http://flyatlas.gla.ac.uk/MidgutAtlas/index.html) for transporter, pump or channel transcripts that show enriched expression in the acid region of the midgut (Supplementary Table 1). Twelve of the thirteen subunits which compose the H\(^+\) V-ATPase are \(\geq2\)-fold up-regulated in the acidic region of the midgut (Fig. 2). The thirteenth subunit (the V\(_o\) ‘a’ subunit), is transcribed from three alternative genes in the larval midgut; \(vha100-2\), \(vha100-4\) and \(vha100-5\) (Fig. 2D–F). These genes show varied expression in the midgut; \(vha100-2\) is expressed throughout but enriched in the acidic region, \(vha100-5\) is expressed throughout but enriched in the transitional region, and \(vha100-4\) is expressed solely in the acidic region. Although it does not provide regional resolution for the midgut, the online atlas of gene expression, FlyAtlas.org\(^\text{20}\), confirms that \(vha100-2\) is expressed throughout larvae and adults, \(vha100-5\) is midgut/hindgut/Malpighian tubule-specific, and \(vha100-4\) is only transcribed in the midgut (Supplementary Fig. 1). The V\(_o\) ‘a’ subunit helps to dock the H\(^+\) V-ATPase complex to the appropriate cellular membrane, and the specificity of \(vha100-4\) transcription suggests it may be required for targeting to the copper cell apical membrane, where it could pump protons into the lumen; V-ATPases invariably pump protons out of the cell, and in insects are commonly concentrated in apical membranes.

![Figure 1. The acidic region of the larval \textit{Drosophila} midgut. (A) Midgut of an intact \textit{Canton S} larva fed \(m\)-Cresol purple pH dye (red pH < 2.4 yellow pH 2.5–8 purple pH > 8); (B) Excised midgut of a \textit{Canton S} larvae maintained on \(m\)-Cresol purple pH dye showing five regions of pH. The anterior acidic region is spatially distant from the posterior alkaline region after midgut dissection (B), but lies parallel in the intact larva (A).](image-url)
plasma membranes, but can also be basally located. With their broader expression profiles (Supplementary Fig. 1), vha100-2 and vha100-5 could play roles in H+ V-ATPase complexes at the basal membrane, or in endomembranes.

To validate the proposed roles for candidate genes, available RNAi stocks for each gene were obtained, and crossed to a Tsp42Ec-Gal4 driver, which drives expression specifically in the caeca, anterior and acid regions of the midgut. In this way, we were able to generate Drosophila larvae with normal gene expression, except for the caeca, anterior and acidic parts of the midgut. These larvae were screened for attenuation of the acid luminal pH with a pH indicator screening assay. The screening assay utilized a range of pH dyes which change colour at a known pH (Fig. 1), allowing us to determine the upper and lower boundary of pH for each midgut region, in each larval genotype tested.

Accordingly, we individually knocked down each vha100 subunit and assessed its effect on acid generation (Fig. 3). The results confirm that the acid region-enriched subunits vha100-2 and vha100-4 each contribute to maximal acidification, but vha100-5 knockdown does not. Using a range of dyes, knockdown of vha100-4 increases pH from pH 2 to pH 4.6–5.2, whereas vha100-2 knockdown results in a more modest increase to pH 3 (Supplementary Table 4).

pH generation requires a network of ion transporters. Analysis of the acidic region RNAseq transcriptome shows enrichment of transcripts involved in proteolysis, metal detoxification, ion transport, and a range of metabolic processes. To investigate the mechanism of pH generation, we identified ion transporters enriched in the acidic region, and assessed the effect of gene knockdown on pH. Five additional genes are involved in pH generation; the potassium/chloride symporter Kcc, the potassium channel Slowpoke, the bicarbonate/chloride exchanger CG8177 and the chloride channel CG11340, together with carbonic anhydrase CAH1 (Fig. 4). Single knockdown of each gene increased pH from 2 units to 3 units (CAH1, CG11340, CG8177), pH 3.1–4.4 (Kcc) or pH 4.6–5.2 (Slowpoke) (Fig. 5, Supplementary Table 4). As in humans, the transport of H+, Cl−, K+ and HCO3− is vital for acid generation in Drosophila. Region-autonomous control of pH is also apparent, as increasing pH in the acidic region does not alter pH in the neighbouring ‘neutral’ region, which is maintained at pH 7.

It is possible to validate some of these findings pharmacologically. Acid generation can also be reduced by feeding larvae Acetazolamide, which blocks carbonic anhydrase function (Fig. 5H). By contrast, as expected, the human H+, K+-ATPase inhibitor Omeprazole did not impede acid pH generation in the Drosophila larval midgut, even at high concentration (35 mg/l in diet, cf. 0.2–1.2 mg/l plasma in humans) (Fig. 5G). Acid generation in insects is thus genetically and pharmacologically distinct from humans.
Figure 3. The H⁺ V-ATPase complex is required for acidic pH generation. Midgut pH was assessed using Thymol blue pH dye (red pH < 2.4, yellow pH 2.5–8, blue pH > 8). Canton S and parental controls (A–C,E,G) all maintain a region of pH 2 (orange/red staining) in the acidic region (annotated by white arrows), whereas acidity is reduced in the vha100-2 (D) and vha100-4 (F) knockdown lines (yellow staining), but not the vha100-5 knockdown line (H).
The acidic region contributes to immune defence. Although maintaining delineated regions of extreme pH is energy-intensive (and thus unlikely to persist unless it confers a selective advantage), disruption of copper cell morphology and function did not affect development or survival in lab-reared Drosophila13,23. What functions could be served by the low pH region? In humans, gut acidity promotes both digestion and defence against infection24. To test whether the acidic region impacts survival when the food source contains potentially hazardous compounds or organisms, first instar Drosophila larvae were subjected to a range of immune and ionic stresses, and their development to pupation assessed. Survival and development of vha100-4 knockdown larvae under optimal rearing was not significantly different from the vha100-4-RNAi and Tsp42Ec-Gal4 parental controls (Fig. 6A), although there is a small but significant decrease in adult life-span (Fig. 6B). To investigate the role of the acidic region in immune defence, mid-L1 larvae were orally infected with Pseudomonas aeruginosa, at a concentration selected to not significantly reduce the survival or development time of wildtype larvae, and monitored to pupation. Knockdown of vha100-4 significantly affected both development time and survival to pupation (Fig. 6C). To determine whether the acidic region impacts the gut bacterial load gained through feeding, L3 larvae were fed Pseudomonas entomophila, marked with a rifampicin expression plasmid, for 2 hours, and the bacterial content of the midgut immediately assessed (Fig. 6D). Although the amounts of food eaten were similar in both cases, Tsp42Ec-Gal4 > vha100-4-RNAi larvae contained ~8-times as many CFUs as the parental controls, suggesting that the acidic region impedes survival or proliferation of pathogenic microbes.

Insects can also be stressed by adverse levels of ions or osmolytes in the diet. Accordingly, survival of vha100-4, slowpoke and kcc knockdown larvae was assessed on exposure to high dietary KCl or NaCl (Fig. 6E, Supplementary Fig. 3). vha100-4 knockdown larvae showed decreased survival upon exposure to both ion loads, as did Slowpoke knockdown larvae. This is reasonable, because the V-ATPase is the primary energizing force in insect epithelia, and drives secondary ion fluxes19. As might be expected, knockdown of the K\(^+\)/Cl\(^-\) cotransporter kazachoc (Kcc) in larvae renders them susceptible to KCl loading, but not NaCl. These data suggest that in addition to generating acid pH, ion transporters up-regulated in the acidic region play an essential role for ion homeostasis in the larva.

Gut pH influences the microbiome, and the microbiome influences gut pH. Given the impact of increased gut pH on Pseudomonas load (Fig. 6D), we investigated whether the pH of the acidic region of the larval Drosophila midgut influences the composition and abundance of the microbiota. Because the gut microbiota of Drosophila can vary, apparently stochastically, even among different cultures of the same genotype25,26, the microbiota in the Drosophila Tsp42Ec-Gal4 > vha100-4 knockdown line and two parental controls was standardized to bacteria of the genera Acetobacter and Lactobacillus (specifically, A. pomorum, A. tropicalis, L. plantarum and L. brevis) that commonly dominate the microbiota of Drosophila in laboratory culture25,27. For each gut region, the cross-sectional area did not vary significantly across the three Drosophila lines (Supplementary Table S2),
Figure 5. Multiple genes are required for pH generation. Midgut pH was assessed using Thymol blue pH dye (red pH < 2.4, yellow pH 2.5–8, blue pH > 8). (A) Control parental RNAi line (acidic region ~pH 2). Knockdown of CAH1 (B), Kcc (C), Slowpoke (D), the SLC4A anion exchanger CG8177 (E) or the ligand-gated chloride channel CG11340 (F) reduced acidity in comparison to parental controls. (G) There was no effect of the H⁺, K⁺-ATPase inhibitor omeprazole (1 mM), but inhibition of carbonic anhydrase activity using the inhibitor compound acetazolamide (100μM) reduced acidity (H).
indicating that gut volumes per region did not differ between the Drosophila lines used; and therefore bacterial abundance was normalized to gut region. In the Tsp42Ec-Gal4 > vha100-4-RNAi larvae, the abundance of Acetobacter was significantly increased in all regions of the midgut distal to the acidic region, relative to the parental lines; and Lactobacillus populations were significantly elevated in the acidic and neutral regions of Tsp42Ec-Gal4 > vha100-4-RNAi larvae (Fig. 7). These data indicate that passage through the extremely low pH in the acidic region of wild-type Drosophila can suppress the populations of bacteria, either in that region or more distally. Similarly, during their study on aging, Li et al. found that commensal bacteria increased in the midgut of adult flies that lacked copper cells due to expression of LabialRNAI.17

Figure 6. *vha100-4* knockdown increases susceptibility to ion loading and *Pseudomonas* infection. (A) Knockdown of *vha100-4* does not impede larval survival to pupation under standard lab conditions, (B) but does result in a significant decrease in adult lifespan. (C) *Pseudomonas* infection significantly increases larval development time and decreases survival to pupation, and (D) Tsp42Ec-Gal4 > vha100-4-RNAi flies have an increased midgut bacterial load after *Pseudomonas* infection. (E) Knockdown of *vha100-4* also compromises larval survival to pupation when maintained on a 2.5% NaCl or 5% KCl diet. Data are expressed as percent pupation (n ~120 larvae) or percent survival (n ~120 flies). Statistically significant differences were assessed by Kaplan-Meier testing with the logrank test (A–C) or one-way ANOVA analysis (D,E), critical level P = 0.05.
We then investigated the converse question: does the presence of the microbiota influence pH in the acidic region? We raised *white* *Drosophila* either axenically (with no gut bacteria) or gnotobiotically with the four bacterial species used previously, and assayed midgut pH in third-instar larvae using thymol blue pH dyes (Supplementary Fig. 4). Gnotobiotic larvae with a microbiota composed of *Acetobacter* and *Lactobacillus* maintained an acidic region of pH 2.0 ± 0.1, (mean ± s.e., N = 10), but the acidic region of the axenic larvae was significantly lower (pH 1.5 ± 0.2, N = 10, t-test P < 0.01). The bacterial cells may thus be able to ameliorate the pH of the acidic region, by the release of weak bases 28. For example, the stomach-inhabiting *Helicobacter pyloris* increases the pH in its immediate environs by ammonia production29. Additionally, the bacteria have a complex relationship with physiological processes which may affect ionic homeostasis in the gut, such as nutrition and immune function30,31, resulting in a change to the pH of the acidic region. Although the underlying mechanisms remain to be established, these experiments demonstrate the importance of the microbiota in shaping the gut internal milieu.

**Conclusions**

This work delineates a set of genes that is necessary to maintain a normal acid region in the midgut, shows that these differ significantly from those implicated in human parietal cell function, and demonstrates that the low pH region serves to control the gut microbiome. The lack of a genomically-encoded K⁺/H⁺ ATPase in *Drosophila*, and corresponding Omeprazole insensitivity that we show here, means that the driving force for acidification must come from elsewhere; as suggested previously13,19–32, the V-ATPase serves such an energizing role in midgut, as in other insect transporting epithelia19–33. By using a new, specific GAL4 driver, we were able to show that expression knockdown of V-ATPase, kazachok, slowpoke or carbonic anhydrase specifically in the midguts of otherwise normal flies, is sufficient to impact the pH of the acidic region.

As in humans, the low pH region of *Drosophila* protects against pathogenic bacterial colonization, and regulates the abundance of non-pathogenic bacteria in the gut. If the low pH region is disrupted, the bacterial populations increase. This communication is two-way, as eliminating the microbiota in turn alters gut pH. Such a balance could help explain why global knockdown of V-ATPase was reported to produce obese flies32, whereas we did not detect such an effect in our midgut-specific V-ATPase knockdowns. There is an increasing awareness that the composition of the gut microbiome in humans can have significant health sequelae34,35; the larval *Drosophila* midgut may provide an excellent and highly tractable system for simple model studies.

**Materials and Methods**

**Drosophila methods.** *Drosophila* were reared on standard diet (10 g agar, 15 g sucrose, 30 g glucose, 35 g dried yeast, 15 g maize meal, 10 g wheat germ, 30 g treacle, 10 g soya flour, per litre)36 in vials, at 26 °C with a 12:12h photoperiod and at 45–55% relative humidity. Where required, they were anesthetized by brief exposure to carbon dioxide. GAL4/UAS crosses were maintained at 26 °C, to ensure strong operation of the GAL4/UAS system. For such experiments, the controls were similarly treated.

**Generation of RNAi alleles.** Commercially available UAS-RNAi stocks were ordered from the Vienna *Drosophila* Research Centre37 and the Transgenic RNAi Project38. Where commercial stocks were unavailable,
RNAi alleles were generated using the pwALIUM20 vector (Transgenic RNAi Project). Inserts were verified by sequencing, and sent for commercial germ-line transformation by BestGene (California, USA), where they were inserted into the AttP2 fly-line using PhiC31 site-specific integration (for primers see Supplementary Table 2).

**Drosophila stocks.** Stocks were either generated or purchased as above. Stocks used were: Canton S (wild-type); AttP2 (background for all P-element insertions used); white^luv^ (for crossing transgenic stocks into the same genetic background), vha100-2-RNAi (VDRC stock 109763), vha100-4-RNAi (generated using pwALIUM20), vha100-5-RNAi (TRIP stock HM04032), CAH1-RNAi (VDRC stock 104429), slowpoke-RNAi (TRIP stock JF01470), kcc-RNAi (TRIP stock HMS01058), CG8177-RNAi (TRIP stock HMC03399), CG11340-RNAi (TRIP stock JF02028), labial-RNAi (TRIP stock JF02317). Tsp42Ec-Gal4 drives expression specifically from the gastric caeca to the border of the acidic and neutral regions of the Drosophila larval midgut. To generate Tsp42Ec-Gal4, the putative promoter sequence of Tsp42Ec was amplified by PCR using Canton S genomic DNA as a template, and cloned into the pStinger vector 39. Inserts were verified by sequencing, sent for commercial germ-line transformation by BestGene (California, USA), and their chromosomes of insertion verified according to standard genetic techniques (for primers see Supplementary Table 2).

**Midgut pH assays.** pH was measured in the larval midgut using Thymol blue, m-Cresol purple, Cresol red, Bromocresol purple, Bromothymol blue, Phenol Red and Congo red indicator dyes (all purchased from Sigma-Aldrich). Indicators were added to melted *Drosophila* diet (0.1% w/v), immediately mixed, and allowed to cool to room temperature. Larvae of the appropriate genotype were added, and after 2 hours the midgut excised in Schneider’s insect medium (Invitrogen). Micrographs were taken immediately using a Sony CyberShot NEX-C3 mounted on a Leica stereo microscope, as pH remains stable for only a few minutes after dissection. Images were processed using Adobe Photoshop CS5.1. Where noted, the diet was also supplemented with 1 mM Omeprazole (Sigma-Aldrich) or 100 μM acetazolamide (Sigma-Aldrich).

**Quantitative RT-PCR.** Gene expression knockdown was assessed in the midgut acidic region of Tsp42Ec-Gal4 > UAS-RNAi larvae, to ensure all RNAi lines had significantly lowered gene expression (see Supplementary Fig. 2). Briefly, dissected midguts were stored in RLT buffer, and RNA extracted using an RNAeasy Mini Kit (Qiagen). PCR was performed in a One-Step real-time PCR machine (Applied Biosystems) using Power SYBR Green RNA-to-CT 1-Step Kit (Applied Biosystems). Expression was quantified using the ΔΔCa method, using α-tubulin as a reference gene, and n = 3 biological replicates 39.

**Larval bacterial infection assays.** First instar larvae of the appropriate genotype were transferred in groups of 30 into a mixture containing 200 μl of mashed banana combined with 200 μl concentrated *Pseudomonas aeruginosa* (TCS Biosciences strain MM41), OD150. Larvae were maintained in the mixture for 30 minutes, and then tipped into a vial containing standard diet and maintained at 26 °C. The vials were scored for pupae morning and evening over 7 days and variation in time to pupation was analyzed by the Kaplan-Meier test. [Supplementary Table 2].

**CFU determination.** For CFU counts, the midguts of 10 larvae (axenic or gnotobiotic) were dissected into each sample to increase accuracy. As a control, the midgut content of larvae fed only on banana was incubated on LB-agar plates for 24 hours under a CO2 atmosphere for 30 minutes, and under aerobic conditions for 24 hours. Images were taken using a WASP-2 instrument, Microbiology International) on mMRS-agar, and incubation under aerobic conditions for *Acetobacter* and under a CO2 atmosphere for *Lactobacillus* 42. The number of CFUs was scored with the Protocol 3 colony counter (Microbiology International). Axenic larvae were assayed as the negative control: none of these homogenates yielded CFUs.

**Statistics.** For normally distributed data with homogenous variance, the significance of differences was assessed with Student’s *t* test, one-way ANOVA or *t* test (two tailed), as appropriate. Where the distribution
shape could not be assumed, Dunn’s multiple comparison post hoc test was used. Significant differences in survival were assessed by testing Kaplan–Meier data with the logrank test. All testing used GraphPad Prism software. Throughout, the critical level is taken as $P = 0.05$.  

References

1. Yao, X. & Forte, J. G. Cell biology of acid secretion by the parietal cell. *Ann Rev Physiol* **65**, 103–131 (2003).
2. Brett, S. Science review: The use of proton pump inhibitors for gastric acid suppression in critical illness. *Critical care* **9**, 45–50 (2005).
3. Miehle, F. E. et al. The cardiac K⁺ channel KCNQ1 is essential for gastric acid secretion. *Gastroenterology* **120**, 1363–1371 (2001).
4. Malinowska, D. H., Kupert, E. Y., Babiniski, A., Sherry, A. M. & Cuppoletti, J. Cloning, functional expression, and characterization of a PKA-activated gastric Cl-channel. *Am J Physiol* **268**, C191–200 (1995).
5. Stuart-Tilley, A., Sardet, C., Pouyssegur, J., Schwartz, M. A., Brown, D. & Alper, S. L. Immunolocalization of anion exchanger AE2 and cation exchanger NHE-1 in distinct adjacent cells of gastric mucosa. *Am J Physiol* **266**, C59–588 (1994).
6. Davenport, H. The inhibition of carbonic anhydrase and of gastric acid secretion by thioacyanate. *Ann J Physiol* **129**, 505–514 (1940).
7. Petrovic, S. et al. Identification of a basolateral Cl/H⁺-ATPase specific to gastric parietal cells. *Am J Physiol Gastrointest Liver Physiol* **284**, G1093–1103 (2003).
8. Wolosin, J. M. & Forte, J. G. Stimulation of oxyntic cell triggers K⁺ and Cl⁻ conductances in apical H⁺⁻K⁺⁻ATPase membrane. *Am J Physiol* **246**, C537–545 (1984).
9. Reenstra, W. W. & Forte, J. G. Characterization of K⁺ and Cl⁻ conductances in apical membrane vesicles from stimulated rabbit oxyntic cells. *Am J Physiol* **259**, G850–858 (1990).
10. Shanbhag, S. & Tripathi, S. Epithelial ultrastructure and cellular mechanisms of acid and base transport in the *Drosophila* midgut. *J Exp Biol* **212**, 1731–1744 (2009).
11. Strand, M. & Micchelli, C. A. Quiescent gastric stem cells maintain the adult *Drosophila* stomach. *Proc Natl Acad Sci USA* **108**, 17696–17701 (2011).
12. Filhie, B. K., Poulton, D. F. & Waterhouse, D. F. Ultrastructure of the copper-accumulating region of the *Drosophila* larval midgut. *Tissue Cell* **3**, 77–102 (1971).
13. Dubreuil, R. R., Frankel, J., Wang, P., Howrylak, J., Kappil, M. & Grushko, T. A. Mutations of alpha spectrin and labial block c-terminal cell differentiation and acid secretion in the middle midgut of *Drosophila* larvae. *Dev Biol.* **194**, 1–11 (1998).
14. Shi, S. C. et al. Drosophila microbiome modulates host developmental and metabolic homeostasis via insulin signaling. *Science* **334**, 670–674 (2011).
15. Lee, W. J. & Brey, P. T. How microbiomes influence metazoan development: insights from history and *Drosophila* modeling of gut-microbe interactions. *Annu Rev Cell Dev Biol* **29**, 571–592 (2013).
16. Lemaitre, B. & Miguel-Aliaia, I. The digestive tract of *Drosophila* melangaster. *Annu Rev Genet* **47**, 377–404 (2013).
17. Li, H., Qi, Y. & Jaspar, H. Preventing age-related decline of gut compartmentalization limits microbiota dysbiosis and extends lifespan. *Cell Host & Microbe* **19**, 240–253 (2016).
18. Buchon, N. et al. Morphological and molecular characterization of adult midgut compartmentalization in *Drosophila*. *Cell Rep* **3**, 1725–1738 (2013).
19. Chintapalli, V. R., Wang, J., Herrzy, P., Davies, S. A. & Dow, J. A. T. Data-mining the FlyAtlas online resource to identify core motifs across transporting epithelia. *BMC Genomics* **14**, 518 (2013).
20. Chintapalli, V. R., Wang, J. & Dow, J. A. T. Using FlyAtlas to identify better *Drosophila* models of human disease. *Nat Genet* **39**, 715–720 (2007).
21. Wiecekorek, H., Gruber, G., Harvey, W. R., Hess, M., Merzendorfer, H. & Zeiske, W. Structure and regulation of insect plasma membrane H⁺⁻V-ATPase. *J Exp Biol* **203**, 127–135 (2000).
22. Wiecekorek, H. & Harvey, W. R. Energization of animal plasma membranes by the proton-motive force. *Physiol Zool* **68**, 15–23 (1995).
23. Dubreuil, R. R., Grushko, T. & Baumann, O. Differential effects of a labial mutation on the development, structure, and function of stomach acid-secreting cells in *Drosophila* melangaster larvae and adults. *Cell Tissue Res* **306**, 167–178 (2001).
24. Giannella, R. A., Broitman, S. A. & Zamechek, N. Influence of gastric acidity on bacterial and parasitic enteric infections. A perspective. *Ann Intern Med* **78**, 271–276 (1973).
25. Chandler, J. A., Lang, I. M., Bhatnagar, S., Eisen, J. I. A. & Kopp, A. Bacterial communities of diverse *Drosophila* species: ecological context of a host-microbe model system. *PLoS Genet* **7**, e1002272 (2011).
26. Wong, A. C., Chaston, J. M. & Douglas, A. E. The inconstant gut microbiota of *Drosophila* species revealed by 16S rRNA gene analysis. *ISME* **7**, 1922–1932 (2013).
27. Wong, C. N., Ng, P. & Douglas, A. E. Low-diversity bacterial community in the gut of the fruitfly *Drosophila* melangaster. *Environ Microbiol* **13**, 1889–1900 (2011).
28. Thomas, K. C., Hynes, S. H. & Inglewed, W. M. Influence of medium buffering capacity on inhibition of Saccharomyces cerevisiae growth by acetic and lactic acids. *Appl Environ Microbiol* **68**, 1616–1623 (2002).
29. Salama, N. R., Hartung, M. L. & Muller, A. Life in the human stomach: persistence strategies of the bacterial pathogen Helicobacter pylori. *Nat Rev Microbiol* **11**, 385–399 (2013).
30. Buchon, N., Silverman, N. & Cherry, S. Immunity in *Drosophila* melangaster—from microbial recognition to whole-organism physiology. *Nat Rev Immunol* **14**, 796–810 (2014).
31. Douglas, A. E. The molecular basis of bacterial-insect symbiosis. *J Mol Biol* **426**, 3830–3837 (2014).
32. Lin, W. S. et al. Reduced Gut Acidity Induces an Obese-Like Phenotype in *Drosophila* melangaster and in Mice. *PLoS One* **10**, e0139722 (2015).
33. Allain, A. K., Du, J., Davies, S. A. & Dow, J. A. Genome-wide survey of V-ATPase genes in *Drosophila* reveals a conserved renal phenotype for lethal alleles. *Physiological genomics* **22**, 128–138 (2005).
34. Clemente, J. C., Urrisell, L. K., Parfrey, L. W. & Knight, R. The impact of the gut microbiota on human health: an integrative view. *Cell 148*, 1258–1270 (2012).
35. Shreiner, A. B., Kao, Y. J. & Young, V. B. The gastrin-gastroenteroendocrine (G) system: a physiologic perspective. *Curr Opin Gastroenterol* **10**, 69–75 (1995).
36. Ashburner, M. *Drosophila: a laboratory manual*. Cold Spring Harbor: Laboratory Press (1989).
37. Dietz, G. et al. A genome-wide transgenic RNAi library for conditional genetic inactivation in *Drosophila*. *Nature* **448**, 151–156 (2007).
38. Ni, J. Q. et al. Vector and parameters for targeted transgenic RNA interference in *Drosophila* melanogaster. *Nat Methods* **5**, 49–51 (2008).
39. Barolo, S., Carver, L. A. & Posakony, J. W. GFP and beta-galactosidase transformation vectors for promoter/enhancer analysis in *Drosophila*. *BioTechniques* **29**, 726, 728, 730, 732 (2000).
40. Stergiopoulos, K., Cabrero, P., Davies, S. A. & Dow, J. A. T. Salty dog, an SLC5 symporter, modulates *Drosophila* response to salt stress. *Physiol Genomics* **37**, 1–11 (2009).
41. Newell, P. D. & Douglas, A. E. Interspecies interactions determine the impact of the gut microbiota on nutrient allocation in *Drosophila* melangaster. *Appl Environ Microbiol* **80**, 788–796 (2014).
42. Newella, P. D. et al. *In vivo* function and comparative genomic analyses of the *Drosophila* gut microbiota identify candidate symbiosis factors. *Front Microbiol* **5**, 576 (2014).
Acknowledgements
We thank the TRiP at Harvard Medical School (NIH/NIGMS R01-GM084947) and VDRC for providing transgenic RNAi fly stocks and plasmid vectors used in this study. This work was funded by the UK Biotechnology and Biological Sciences Research Council (to JATD and SAD) and NSF grant BIO 1241099 (to AED).

Author Contributions
J.A.T.D. and S.A.D. obtained grant funding. G.O., J.A.T.D., S.A.D. and A.E.D. designed the experiments. G.O., L.H. and Y.L. performed the experiments. G.O., J.A.T.D. and A.E.D. wrote the manuscript text. G.O. and Y.L. prepared the figures. All authors reviewed the manuscript.

Additional Information
Supplementary information accompanies this paper at http://www.nature.com/srep

Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Overend, G. et al. Molecular mechanism and functional significance of acid generation in the Drosophila midgut. Sci. Rep. 6, 27242; doi: 10.1038/srep27242 (2016).

This work is licensed under a Creative Commons Attribution 4.0 International License. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/