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A Genome-wide Screen Identifies PAPP-AA-Mediated IGFR Signaling as a Novel Regulator of **Habituation Learning**

Highlights

- Genome-wide genetic screen identifies first set of vertebrate learning mutants
- pregnancy-associated plasma protein-aa is a novel regulator of habituation learning
- PAPP-AA functions as a metalloprotease to promote habituation
- PAPP-AA/IGF1R/PI3K/Akt signaling axis as a pharmacological target for learning

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In Brief

Through a forward genetic screen in zebrafish, Wolman et al. isolate the first functional vertebrate gene set for habituation learning and identify pregnancy-associated plasma protein aa (PAPP-AA)-regulated IGF signaling as a novel mechanism underlying habituation learning.



A Genome-wide Screen Identifies PAPP-AA-Mediated IGFR Signaling as a Novel Regulator of Habituation Learning

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SUMMARY

Habituation represents a fundamental form of learning, yet the underlying molecular genetic mechanisms are not well defined. Here we report on a genome-wide genetic screen, coupled with wholegenome sequencing, that identified 14 zebrafish startle habituation mutants including mutants of the vertebrate-specific gene pregnancy-associated plasma protein-aa (pappaa). PAPP-AA encodes an extracellular metalloprotease known to increase IGF bioavailability, thereby enhancing IGF receptor signaling. We find that *pappaa* is expressed by startle circuit neurons, and expression of wild-type but not a metalloprotease-inactive version of pappaa restores habituation in pappaa mutants. Furthermore, acutely inhibiting IGF1R function in wild-type reduces habituation, while activation of IGF1R downstream effectors in pappaa mutants restores habituation, demonstrating that pappaa promotes learning by acutely and locally increasing IGF bioavailability. In sum, our results define the first functional gene set for habituation learning in a vertebrate and identify PAPPAA-regulated IGF signaling as a novel mechanism regulating habituation learning.

INTRODUCTION

All animals exploit a fundamental mechanism of non-associative learning, called habituation, to filter irrelevant input and prioritize attention (Thompson and Spencer, 1966). Habituation is characterized by progressive response decline to repeatedly experienced yet inconsequential stimuli (Groves and Thompson, 1970; Rankin et al., 2009). Despite its simplicity, habituation is an attractive form of learning because it provides a quantifiable form of neuroplasticity (Poon and Young, 2006). Deficits in habituation represent hallmark features of human cognitive and behavioral disorders, including schizophrenia, addiction, attention deficit hyperactivity disorder, and other disorders marked by "intellectual disability" (Braff et al., 1992; Jansiewicz et al., 2004; McSweeney et al., 2005). Despite its biological conservation and clinical relevance, our understanding of the genetic mechanisms governing habituation is limited. Identifying the genetic program that governs how neural circuits regulate habituation is therefore instrumental to understanding disorders marked by habituation deficits and for dissecting the genetic basis of higher cognition.

To identify the genetic program that governs vertebrate habituation learning, we took an unbiased, genome-wide approach to define a core set of genes critical for habituation of the vertebrate acoustic startle response. Inspired by behavioral screens in Drosophila and C. elegans (Benzer, 1967; Brenner, 1974; Eddison et al., 2012; Ikeda et al., 2008; L'Etoile et al., 2002; Lau et al., 2013; Pierce-Shimomura et al., 2008; Rankin, 2004; Rankin et al., 1990; Swierczek et al., 2011; Wolf et al., 2007), we performed a forward genetic screen using a high-throughput behavior testing apparatus that measures zebrafish startle habituation (Wolman et al., 2011) and then applied whole-genome sequence (WGS) analysis to molecularly identify the mutated genes. Here, we report on (1) a set of 14 mutants with specific deficits in startle habituation, (2) the molecular identification of two mutants, and (3) the characterization of a novel and vertebrate specific modulator of zebrafish habituation learning.

As predicted, our gene set includes genes with "expected" functions and genes previously not associated with habituation learning. Specifically, we identified mutations in the *pyruvate carboxylase a (pcxa)* gene, which encodes a rate limiting enzyme in the production of the glutamate (Hertz et al., 2007), a key neurotransmitter for habituation learning (Bespalov et al., 2007; Bickel et al., 2008; Riedel et al., 2003; Rose and Rankin, 2006). Conversely, we also identified a mutation in the vertebrate specific gene *pregnancy associated plasma protein-aa (pappaa)*. PAPP-AA has not been implicated in any type of learning and is known to act as an extracellular metalloprotease to enhance local insulin-like growth factor (IGF) signaling by cleaving IGF binding protein 4 (IGFBP4), which normally restricts IGF from signaling through cell-surface IGF receptors (Conover et al.,





2004; Laursen et al., 2001, 2007; Lawrence et al., 1999). Using a combination of molecular-genetic, pharmacological, and behavioral analyses, we uncover a previously unknown role for PAPP-AA-regulated IGF signaling in mediating habituation learning through an acute and presumptive local mechanism. Overall, our results define the first in vivo function-based set of genes regulating vertebrate habituation learning.

RESULTS

Forward Genetic Screen Identifies Zebrafish Mutants with a Startle Habituation Deficit

By 5 days of age, zebrafish larvae perform a repertoire of simple sensorimotor behaviors that operate on characterized and accessible neural circuits (Wolman and Granato, 2012). For example, exposure to abrupt acoustic stimuli elicits a startle response, an evolutionary conserved and stereotyped yet Figure 1. Genetic Screen Identifies Mutations Affecting Acoustic Startle Habituation (A) Schematic of acoustic startle habituation assay. Larvae are exposed to 10 non-habituating acoustic stimuli, delivered at 20 s interstimulus intervals (ISI), and then 30 habituating stimuli at a 1 s ISI.

(B) Mean acoustic startle habituation percentage calculated by comparing the average frequency of startle responsiveness of an individual to stimuli 1–10 and stimuli 31–40 (Wolman et al., 2011). Behaviorally defined wild-type siblings shown in white bars, mutants in gray bars.

(C) Estimated truncated PCXA^{p171} protein in *information overload*^{p171} mutants due to Y1163X mutation. BC, biotin carboxylase; CT, carboxyl transferase; PT, pyruvate carboxylase tetramerization; BCCP, biotin-carboxy carrier protein.

(D) Estimated truncated PAPP-AA^{p170} protein product in *unfiltered*^{p170} mutants due to Q322X mutation. LG, laminin G-like module; LNR, Lin-12/ Notch repeats, MPD, metzincin proteolytic domain containing zinc-binding consensus sequence (Zn) and Met-turn motif (M); CCP, complement control protein modules 1–5.

E) pappaa^{p170} larvae injected with increasing doses of wild-type pappaa *mRNA* show improved habituation at 5 dpf. *p < 0.01, **p < 0.001, ANOVA with Bonferonni correction versus wild-type sibling (B) or uninjected pappaa^{p170} (E) larvae. N = number of larvae shown within or below each bar. Error bars indicate SEM.

modifiable behavior (Burgess and Granato, 2007b; Eaton et al., 1977; Kimmel et al., 1974; Wolman et al., 2011). Repeated acoustic stimulation rapidly prompts habituation by the larvae with identical kinematic and pharmacodynamic parameters observed in adult zebrafish and mammals (Bespalov et al., 2007; Bickel et al., 2008; Riedel et al., 2003; Wolman et al., 2011). To identify

genes critical for habituation, we mutagenized adult males using ENU and implemented a three-generation breeding scheme to generate homozygous mutant larvae (Dosch et al., 2004; Mullins et al., 1994). For each F3 clutch, we tested 32 larvae at 5 days post-fertilization (dpf) for short-term habituation to repetitive acoustic stimuli (Wolman et al., 2011). Larvae with morphological defects, hearing loss, or aberration in the highly stereotyped kinematics of the startle response were excluded from subsequent analyses. Heritability of the genetic lesion was verified by observing similarly reduced habituation in subsequent generations.

To identify mutants with habituation defects, we used a highthroughput behavioral platform that measures habituation of the acoustic startle response (Wolman et al., 2011). Specifically, 5 dpf larvae were first exposed to ten acoustic stimuli separated by a 20 s interstimulus interval (ISI) to determine baseline startle responsiveness, and then were given 30 acoustic stimuli with a 1 s ISI to evaluate habituation (Figure 1A). Wild-type larvae

Table 1. Zebrafish Habituation Mutants						
Mutant Allele	Startle Habituation	Visual Habituation Percentage	Baseline Activity	Startle Kinematics	Startle Sensitivity	Gene Locus
unfiltered ^{p170}	4.9% ± 1.2%*	No response ^a	Normal	Normal	Increased	рарраа
information overload ^{p171}	23.6% ± 1.6%*	Normal	Normal	Normal	Normal	рсха
ignorance isbliss ^{p172}	17.4% ± 2.3%*	Normal	Reduced	Normal	Increased	Ch. 15 z4396; z9189z13822
irresistible ^{p173}	18.8% ± 2.1%*	Normal	Normal	Normal	Increased	Ch. 7 <i>z7958</i>
slow learner ^{p174}	7.9% ± 2.0%*	Reduced	Reduced	Normal	Increased	Ch. 4 <i>z1366; z1525z4951; z7104</i>
uninhibited ^{p175}	15.7% ± 2.1%*	Normal	Normal	Normal	Increased	n.d.
groundhog day ^{p176}	20.1% ± 1.9%*	Normal	Normal	Normal	Increased	n.d.
dory ^{p177}	13.8% ± 1.9%*	Normal	Reduced	Normal	Increased	n.d.
divided attention ^{p178}	$24.0\% \pm 2.4\%^*$	Normal	Normal	Normal	Increased	n.d.
oops I did itagain ^{p179}	16.6% ± 2.3%*	Reduced	Reduced	Normal	Normal	n.d.
repeat offender ^{p180}	20.5% ± 2.0%*	n.d.	Normal	Normal	Increased	n.d.
fool me twice ^{p181}	21.49% ± 4.48%*	n.d.	n.d.	Normal	Increased	n.d.
forgetful ^{p182} ;forgetful ^{p183}	28.49% ± 3.1%*	n.d.	n.d.	Normal	Normal	n.d.

Summary of behavioral analyses, including acoustic startle habituation (Figure 1B), visual habituation (Figure S2C), baseline activity (Figure S2B), startle kinematics, and startle sensitivity (Figure S2A) of mutants versus wild-type siblings. Startle kinematic analysis included latency to initiate C-Bend, turning angle during Cbend, and duration of C-bend (primary data not shown). Genetic locus column shows gene or chromosomal region linked to mutant phenotype. Results from complementation crosses suggest these 14 mutants represent alleles of 13 genes. *p < 0.01 mutants versus wild type siblings using ANOVA with Bonferonni correction. n.d., not determined.

^aNo response indicates that *pappaa^{p170}* mutants do not perform the stereotyped O-bend maneuver to the "dark flash" stimuli and therefore cannot be evaluated for visual habituation.

show a rapid reduction in startle response initiation and stereotypically habituate by more than 80% under these conditions (Wolman et al., 2011). Therefore, clutches with approximately 15%–25% of the larvae habituating by less than 50% indicated that the larvae were homozygous for a recessive mutation affecting habituation. Larvae habituating by less than 50% were classified "mutant" (Figure 1B; Movie S1). Using this approach, we screened 405 mutagenized F2 families, corresponding to 614 genomes, and identified 14 habituation mutants (Table 1; Figure 1B).

The behavioral severity and specificity of each of the 14 mutants was determined by comparing the magnitude of the habituation deficit and performing additional behavioral analyses, including acoustic startle sensitivity, baseline activity level, and habituation to repetitive visual stimuli. A comparative analysis revealed that startle habituation in the 14 mutants was affected to different degrees, ranging from 24% habituation (information overload^{p171}, divided attention^{p178}) down to the almost complete absence of habituation (unfiltered^{p170}; Table 1; Figure 1B). We further evaluated two of the mutants. unfiltered^{p170} and information overload^{p171}, for short-term habituation to acoustic stimuli delivered at longer ISIs and found that both mutants also show strong habituation deficits to the less frequently delivered stimuli (Figure S1). Startle sensitivity was significantly enhanced in unfiltered^{p170}, ignorance is bliss^{p172}, irresistible^{p173}, slow learner^{p174}, uninhibited^{p175}, groundhog day^{p176}, dory^{p177}, divided attention^{p178}, repeat offender^{p180}, and fool me twice^{p181}, but not information overload^{p171}, oops I did it again^{p179}, or forgetful^{p182,p183} mutants (Table 1; Figure S2A). Recording gross movement of unstimulated larvae revealed that none of the mutants displayed

spontaneous hyperactivity (Table 1; Figure S2B). In fact, ignorance is bliss^{p172}, slow learner^{p174}, dory^{p177}, and oops I did it again^{p179} mutants performed less spontaneous, overall movement compared to their siblings (Table 1: Figure S2B). Larvae were also given a series of repetitive visual "dark flashes" (Burgess and Granato, 2007a) to determine whether the acoustic startle habituation mutants were capable of habituating to visual stimuli (see Experimental Procedures). Both slow learner^{p174} and oops I did it again^{p179} mutants showed visual habituation deficits, suggesting circuit and/or molecular overlap between acoustic startle and visual habituation (Table 1; Figure S2C). Thus, a genome-wide genetic screen identified a set of mutants in which acoustic startle habituation is reduced to varying degrees. None of these mutants display obvious deficits in acoustic startle performance (i.e., kinematic parameters) or exhibit increased baseline activity, further underscoring the specificity of our screening assay. Several mutants exhibit deficits in acoustic stimulus sensitivity or in visual habituation, reflecting potential overlap between the neural circuitry and genetic programs underlying these behaviors and startle habituation.

WGS Identifies Novel Regulators of Habituation Learning

To determine the molecular identity of the mutated genes, we selected two mutants with varying degrees of habituation capacity and differing behavioral profiles and performed WGS. WGS analysis from behaviorally defined *unfiltered*^{p170} and *information overload*^{p171} mutant larvae followed by homozygosity analysis (see Experimental Procedures; Figure S3) identified distinct chromosomal intervals for each of these mutants, which we

confirmed using bulk segregant analysis (Jain et al., 2011). Within these chromosomal intervals, sequencing data revealed unique nonsense mutations in *pyruvate carboxylase a (pcxa)* in *information overload*^{p171} mutants, and in *pregnancy-associated plasma protein-aa (pappaa)* in *unfiltered*^{p170} mutants, respectively (Figures 1C and 1D).

48 hpf

pLL

pappaa

all

A'

С

Α

В

PCXA is a biotin-dependent, mitochondrial enzyme that catalyzes the carboxylation of pyruvate to oxaloacetate, a critical step in the synthesis of glucose, fat, amino acids, and neurotransmitters, including glutamate, which is known to influence habituation (Bespalov et al., 2007; Bickel et al., 2008; Jitrapakdee et al., 2008; Rankin and Wicks, 2000; Riedel et al., 2003). Sequencing of pcxa cDNA from behaviorally identified information overload^{p171} mutant larvae identified a single nucleotide nonsense mutation in pcxa (nt3489: C to A), which causes a premature stop codon within the biotin-carboxy carrier protein (BCCP) domain (Figure 1C). Mutations to the BCCP domain are associated with the most severe form of pyruvate carboxylase deficiency (type B) due to loss of enzymatic activity (Monnot et al., 2009). Future experiments will determine whether the pcxa^{p171} allele shows reduced enzymatic activity as a result of reduced biotin binding and/or catalytic activity due to improper presentation of biotin to the biotin carboxylase (BC) and/or carboxyl transferase (CT) domains.

Sequencing of *pappaa* cDNA from behaviorally identified *unfiltered*^{p170} larvae confirmed a single nucleotide nonsense mutation in *pappaa* (nt964: C to T). This mutation causes a premature stop codon in exon 3 at amino acid 322 of 1,591, severely truncating PAPP-AA upstream of the metzincin proteolytic domain required for substrate proteolysis and upstream of the C-terminal domains required for membrane tethering via heparin sulfate proteoglycans (Figure 1D) (Boldt et al., 2001; Laursen et al., 2002; Weyer et al., 2004). To confirm that mutations in the *pappaa* gene cause the habituation deficit that we observe in *pappaa/unfiltered*^{p170} mutants, we injected wild-type zebrafish *pappaa mRNA* into one-cell stage *pappaa*^{p170} embryos. This restored startle habituation in a dose-dependent

manner, demonstrating that the truncating mutation in zebrafish *pappaa* causes the habituation deficit (Figure 1E). Thus, WGS analysis reveals *pappaa* as a novel regulator of habituation learning.

5 dpf

Mauthr

pappaa Is Expressed throughout the Acoustic Startle Circuit

To understand how pappaa modulates startle habituation, we first examined the spatial expression pattern of pappaa mRNA. The acoustic startle response is triggered by activation of one of the bilateral pair of Mauthner neurons in the hindbrain (Figures 2A and 2A'). Mauthner neurons receive acoustic inputs from the ear (via the statoacoustic ganglia) and the lateral line and send descending commissural axons down the spinal cord to stimulate contralaterally positioned motor neurons and induce contralateral trunk muscle contraction (Eaton and Emberley, 1991; Eaton et al., 2001; Faber et al., 1989; Faber and Korn, 1978; Liu and Fetcho, 1999). Wholemount in situ hybridization revealed that during embryonic development pappaa mRNA is detectable in several sensory components of the startle circuit: in neurons of the anterior and posterior lateral line ganglia and in neurons of the statoacoustic ganglia, which receive afferent input from sensory hair cells in the lateral line neuromasts and the inner ear, respectively (Figures 2A and 2B). Importantly, at 5 dpf, when pappaa^{p170} mutants display habituation defects, pappaa mRNA is detectable in the startle command neurons, the Mauthner neurons (Figure 2C). pappaa mRNA is also expressed by several clusters of neighboring hindbrain interneurons known to modulate Mauthner activation (Figure 2C), including passive hyperpolarizing (PHP) neurons, spiral fiber neurons, and the feedback inhibitory neurons (Faber et al., 1989; Hackett and Faber, 1983; Koyama et al., 2011; Lorent et al., 2001; Scott et al., 1994). Molecular markers for these cell types are currently unavailable yet the location of pappaa mRNA-positive hindbrain neurons is consistent with the location of PHP, spiral fiber, and feedback inhibitory neurons

Figure 2. *pappaa* Expression in Neurons of the Acoustic Startle Circuit

(A and A') Schematic representation of acoustic startle circuit at larval stage from a lateral (A) and dorsal (A') perspective. The acoustic startle circuit includes cranial ganglion (blue), Mauthner neuron and homologs (green), spiral fiber neurons (orange), passive hyperpolarizing (PHP) neurons (red), and feedback inhibitory neurons (purple).

(B and C) In situ hybridization for *pappaa* at 48 hpf (B, purple) and 5 dpf (C, red). Brackets (A and B) mark site of hindbrain neurons controlling startle behavior. (C) Dorsal view, anterior to the top. *pappaa mRNA* in red, Mauthner neuron (M) in green. Arrowheads mark site of spiral fiber neurons, asterisks mark position of PHP neurons, and arrows indicate location of feedback inhibitory neurons. SAG, statoacoustic ganglion; aLL, anterior lateral line ganglion; M, Mauthner. Scale bars = 50 μ m (B) and 10 μ m (C).



Figure 3. Acoustic Startle Circuit Appears Intact in *pappaa*^{p170} Larvae

(A–C) DASPEI labeling (A and B) and mean number (C) of sensory neuromasts.

(D–I and K–L) Projections of confocal stacks acquired at 5 dpf. Lateral views, anterior to the left in (A), (B), (D), and (E). Dorsal views, anterior to the top in (F)–(I) and (K)–(L). (D and E) Arrowheads mark projections (anti-neurofilament, red) from cranial ganglia (anti-HuC, green) to hindbrain. (F–I, K, L) Green label marks *hspGFF130DMCA:UASgap43-citrine* expression in Mauthner and Mid2cm Mauthner homologs. (F and G) Open arrows mark Mauthner (M) axon cap. Open arrowhead marks spiral fiber neuron projection and closed arrows mark contralateral passive hyperpolarizing (PHP) projection. (H and I) Brackets mark lateral dendrite of Mauthner with Cx35-positive club endings. Asterisk marks blood cell.

(J) Mean number of club endings.

(K and L) Glycineric receptors on Mauthner soma. SAG, statoacoustic ganglion; aLL, anterior lateral line ganglion; pLL, posterior lateral line ganglion. N, larvae shown within bars. Error bars indicate SEM. Scale bars = 1 mm (B), 50 μ m (E), and 10 μ m (G, I, and L).

morphology between $pappaa^{p170}$ mutants and siblings (Movie S2 $(pappaa^{+/+}, n = 16)$ and Movie S3 $(pappaa^{p170}, n = 14)$; Figure S4). Similarly, large synaptic club endings on the lateral dendrite, the sites of acoustic input to the Mauthner, and glycinergic receptor distribution on the soma of Mauthner neurons revealed no differences when compared to wild-type animals (Figures 3H–3L; n = $12 pappaa^{p170}$, $pappaa^{+/+}$ n = 10). Taken together, in $pappaa^{p170}$ mutant larvae, sensory afferents to the Mauthner neurons, the Mauthner neurons, and modula-

(Koyama et al., 2011). Thus, *pappaa* is expressed by identified hindbrain neurons that are well characterized components of the acoustic startle circuit.

Neuronal Development of the Acoustic Startle Circuit Is Intact in *pappaa^{p170}* Mutants

Given the expression of *pappaa* in several neuronal cell types known to regulate startle modulation and/or execution, we used vital dye and immunolabeling to examine whether *pappaa* function is required for the development and/or cellular and synaptic integrity of startle circuitry neurons. Analysis of sensory neuromasts that detect acoustic stimuli, sensory cranial ganglia, Mauthner neurons, and some of their inputs from other hindbrain modulatory interneurons such as PHP and spiral fibers did not reveal any obvious defects in 5 dpf *pappaa*^{*p*170} mutants when compared to wild-type animals (Figures 3A–3G; n = 22 pappaa^{*p*170}, n = 24 pappaa^{*p*170}). Furthermore, high-resolution imaging of the Mauthner neuron showed indistinguishable cellular

tory inputs from neighboring hindbrain neurons appear morphologically indistinguishable from those in wild-type.

PAPP-AA Regulates Habituation Learning through Its Metalloprotease Activity

At 5 dpf, when $pappaa^{p_{170}}$ larvae first become behaviorally distinct from their siblings, they appear grossly normal but often fail to fully inflate their swim bladder (62%, n = 26 $pappaa^{p_{170}}$; Figure 4A). By 9 dpf, the $pappaa^{p_{170}}$ mutants are noticeably smaller in size than their wild-type siblings (3.93 mm ± 0.07, n = 15 $pappaa^{p_{170}}$ versus 4.65 mm ± 0.03 SEM, n = 24 $pappaa^{+/+}$). Behavioral testing of $pappaa^{p_{170}}$ mutants at 12 dpf revealed a clear deficit in habituation learning, strongly suggesting that PAPP-AA also mediates habituation learning during post-developmental stages (Figure 4B). $pappaa^{p_{170}}$ mutants die at 16.9 dpf (±0.29 days SEM, n = 10; Figure 4C), precluding the analysis of adult mutants. Therefore, we conducted all future experiments between 5 and 12 dpf.





Figure 4. *pappaa^{p170}* Mutants Show Reduced Swim Bladder Inflation, Habituation, and Survival

(A) *pappaa^{o170}* mutants appear grossly normal at 5 dpf, with exception of some showing an uninflated swim bladder.

(B) pappaa^{p170} larvae show startle habituation deficit at 5 and 12 dpf.

(C) Kaplan-Meier survival curve shows reduced viability in *pappaa*^{p170} mutants. *p < 0.001, one-way ANOVA (with Bonferonni correction) versus pappaa^{+/+}, **p < 0.0001 Mantel-Cox test versus wild-type siblings. N = number of larvae shown within bars (B) or in legend (C). Error bars indicate SEM.

PAPP-A encodes an extracellular metalloprotease known to cleave IGF binding protein 4 (IGFBP4) and hereby increasing insulin-like growth factor (IGF) locally available to bind its receptor (Conover et al., 2004; Laursen et al., 2007; Laursen et al.,



Figure 5. PAPP-AA Metalloprotease Activity Is Required for Startle Habituation

Mean habituation percentage after injection of human wild-type *pappa mRNA* or proteolytically inactive *pappa^{E483A} mRNA*. *p < 0.001, ANOVA with Bonferonni correction versus DMSO treated larvae of the same genotype. N, larvae shown within bars. Error bars indicate SEM.

2001; Lawrence et al., 1999). To ask whether PAPP-AA functions as a protease for habituation learning, we tested the ability of a previously characterized version of human PAPP-A lacking protease activity (*h*-*pappa*^{E483A}) to restore habituation learning in zebrafish *pappaa*^{p170} mutant larvae (Boldt et al., 2001). While control injections of *h*-*pappa mRNA* into one-cell stage embryos significantly improved habituation in *pappaa*^{p170} mutants, injection of equimolar amounts of *h*-*pappa*^{E483A} *mRNA* did not improve habituation deficits in *pappaa*^{p170} mutants (Figure 5). Importantly, RT-PCR analysis revealed that wild-type *h*-*pappa* and *h*-*pappa*^{E483A} *mRNA* persisted through 5 dpf (Figure S5), demonstrating that PAPP-AA metalloprotease activity is required for acoustic habituation learning.

PAPP-AA Regulates Habituation through Acute Control of IGF1R Signaling

Given that PAPP-AA's metalloprotease activity is required for habituation learning, we next asked whether PAPP-AA regulates this process through canonical IGFR signaling. For this we exposed pappaa^{p170} mutants to SC79, a pharmacological activator of Akt, a canonical downstream effector of IGF1R signaling (Anlar et al., 1999; Jo et al., 2012; Laviola et al., 2007). SC79 treatment from 1 to 5 dpf improved habituation in pappaa^{p170} mutants to near wild-type levels (Figure 6A), suggesting that PAPP-AA regulates acoustic startle habituation through IGFR1-Akt signaling. We next asked whether PAPP-AA acts throughout the period of startle circuit development, or whether PAPP-AA regulates IGF1R-Akt signaling acutely during the habituation process. We hypothesized that periods of SC79 exposure sufficient to improve habituation in pappaap¹⁷⁰ mutants would indicate critical periods of PAPP-AA/ IGF1R/Akt signaling. SC79 treatment restricted prior to 3 dpf failed to improve habituation in pappaa^{p170} mutants. In contrast, SC79 treatment between 3 and 5 dpf, even beginning as late as 5 dpf, restored habituation in pappaa^{p170} mutants. Consistent with these results, acute exposure to 740 Y-P, a cell-permeable phosphopeptide activator of PI 3-kinase (PI3K) (Williams and



Figure 6. PAPP-AA Mediates Habituation through Acute Regulation of IGF Signaling

(A) Mean acoustic startle habituation percentage at 5 dpf of larvae treated with DMSO or 1 μ M SC79. Treatment period (dpf) indicated below bars.

(B) Mean acoustic startle habituation percentage of 5 dpf larvae treated with 740 Y-P for 30 min prior to and during habituation assay.

(C and D) Mean acoustic startle habituation percentage after 30 min treatment with BMS-754807 or DMSO at 5 dpf (C) or 12 dpf (D). *p < 0.01, **p < 0.001, ANOVA with Bonferonni correction versus DMSO-treated larvae of the same genotype. N = number of larvae shown within bars. Error bars indicate SEM.

Doherty, 1999), also improved habituation in *pappaa^{p170}* mutants (Figure 6B). Finally, exposure to BMS-754807, a pharmacological inhibitor of IGF1R (Carboni et al., 2009; Kamei et al., 2011) for 30 min prior to and during habituation testing, also significantly reduced startle habituation in a dose-dependent manner in wild-type zebrafish at both 5 and 12 dpf (Figures 6C and 6D). Together, these results reveal a novel, acute role for PAPP-AA/IGF1R/PI3K/Akt signaling in promoting startle habituation.

DISCUSSION

Habituation is a fundamental form of learning during which an animal's response to repetitive, identical stimuli gradually declines (Groves and Thompson, 1970; Thompson and Spencer, 1966). This decline is not due to sensory adaptation or motor fatigue (Rankin et al., 2009; Thompson and Spencer, 1966; Wolman et al., 2011) and is conserved across species. In addition to its conservation, habituation is a particularly interesting form of learning because it provides a measurable form of neuroplasticity that enables animals to ignore irrelevant stimuli in favor of higher-priority stimuli (Poon and Young, 2006), and in humans disruption of habituation is strongly correlated with cognitive impairments (Braff et al., 1992; Jansiewicz et al., 2004; McSweeney et al., 2005). Despite extensive characterization of various forms of habituation learning, including olfactory, mechanosensory, and startle habituation (Engel and Wu, 2009; Giles and Rankin, 2009; Glanzman, 2009; Halberstadt and Geyer, 2009; Koch and Schnitzler, 1997; Schmid et al., 2003; Wilson, 2009), there remains a clear need for functional gene sets underlying habituation learning, particularly in vertebrates. Here, we report on results from a genome-wide genetic screen to identify a gene set selected solely for a functional role in startle habituation. Importantly, this work identifies pappaa as a novel, vertebratespecific regulator of startle habituation. Based on a series of in vivo experiments, we propose a model by which PAPP-AA acts as a metalloprotease to promote startle habituation by increasing IGF receptor signaling during the process of habituation (Figure 7).

A Forward Genetic Screen Identifies Startle Habituation Mutants

With the exception of an unbiased, forward genetic screen for olfactory habituation mutants in Drosophila (Eddison et al., 2012; Wolf et al., 2007), only candidate gene approaches have been used to identify and dissect genetic mechanisms underlying habituation (Castro-Alamancos and Torres-Aleman, 1994; Das et al., 2011; Engel and Wu, 1998; Lau et al., 2013; Morrison and van der Kooy, 2001; Ohta et al., 2014; Rankin and Wicks, 2000; Rose et al., 2003; Sanyal et al., 2004; Swierczek et al., 2011; Typlt et al., 2013). To complement previous work and identify a broad set of core genes critical for vertebrate startle habituation, we conducted a classical three generation forward genetic screen. This approach yielded a set of 14 mutants with unaltered startle performance but impaired startle habituation (Figure 1). Because these mutants do not exhibit any morphological or motility defects, they probably represent mutants previously not identified in the major zebrafish screens from Tubingen





PAPP-AA cleaves IGFBP to increase bioavailable IGF. "Free" IGF then binds and signals through the IGF1R, which triggers activation of PI3K and Akt to regulate habituation.

and Boston in the mid-1990s (see Development, 1996, Issue 123). We noticed that several of the mutants displayed enhanced sensitivity to acoustic stimuli (Figure S2A), consistent with the idea that the mechanisms controlling startle sensitivity are located "upstream" within the neural circuitry regulating startle habituation (Hoffman and Searle, 1968; Pilz and Schnitzler, 1996). Importantly, not all habituation mutants exhibit increased startle sensitivity (e.g., information overload^{p171}, oops I did it again^{p179}, and forgetful^{p182,p183}). Conversely, in a companion screen, we identified mutants with much greater enhanced startle sensitivity than the mutants described here, yet these mutants habituate normally (data not shown). We find that several habituation mutants (e.g., information overload^{p171}, ignorance is bliss^{p172}, irresistible^{p173}, slow learner^{p174}, uninhibited^{p175}, groundhog day^{p176}, and divided attention^{p177}) are homozygous viable and display startle habituation deficits when tested as young adults (~3 months), confirming that the molecular-genetic mechanisms driving habituation learning in 5 dpf larvae persists into adulthood. Finally, we find that two of the mutants with reduced habituation to acoustic stimuli also displayed visual habituation deficits (Figure S2C; slow learner⁰¹⁷⁴ and oops I did it again^{p179}), providing evidence that these two affected genes play a role in habituation independent of sensory modality. Thus, we have identified the first set of startle habituation genes characterized solely by their functional requirement for habituation, and given that we previously demonstrated that acoustic startle habituation in zebrafish larvae can be reversed, i.e., dishabituated by a novel stimulus (Wolman et al., 2011), we expect to find already well-characterized genes as well as genes previously not known to affect the central process of habituation learning.

A Role for *pyruvate carboxylase a* in Habituation Learning

Whole-genome sequence analysis and high-resolution linkage analyses reveal that the information overload^{p171} habituation phenotype co-segregates with a premature stop codon in the biotin-carboxy carrier protein (BCCP) domain of the pyruvate carboxylase a (pcxa) gene (Figure 1C). PCXA catalyzes the carboxylation of pyruvate to oxaloacetate, a necessary step toward the production of glucose, fat, and amino acids, including glutamate (Jitrapakdee et al., 2008). Importantly, pyruvate carboxylase activity is undetectable in neurons but is active in astrocytes, where it plays a pivotal role in the maintenance of the glutamate neurotransmitter pool (Hertz, 2004; Schousboe et al., 2013; Whitfield et al., 1996). In humans, pyruvate carboxylase deficiency results in severe psychomotor retardation and affected individuals die within months to a few years after birth (Monnot et al., 2009). It might therefore appear surprising that pcxa^{p171} mutant zebrafish larvae appear grossly normal and that they are at least partially homozygous viable (data not shown). One likely explanation for the viability of pcxa mutants is that a second paralog gene, pyruvate carboxylase b (pcxb) located on chromosome 7, attenuates the impact of pcxa deficiency. How then does pcxa affect habituation learning? It is well documented that glutamate neurotransmission regulates habituation learning in both mammals and zebrafish (Bespalov et al., 2007; Bickel et al., 2008; Roberts et al., 2013; Wolman et al., 2011). It is therefore tempting to speculate that glia associated with neurons of the startle circuit require pcxa to maintain a constant pool of glutamate. Further analyses are necessary to test this hypothesis and to understand how pyruvate carboxylase-dependent glutamate synthesis promotes neuronal plasticity, including habituation learning.

PAPP-AA Is a Novel Regulator of Habituation Learning and Acutely Promotes IGFR Signaling

Mammalian pregnancy-associated plasma protein a (PAPP-A) was originally purified from late pregnancy plasma and later shown to encode a membrane associated metalloprotease that specifically cleaves insulin-like growth factor binding proteins IGFBP-4 and IGFBP-5, thereby releasing IGF from their binding partners and hence promoting IGF receptor activation (Laursen et al., 2001, 2007; Lawrence et al., 1999; Lin et al., 1974). In mice, knockout of pappa results in viable offspring about 60% the size of wild-type at birth, identical to the phenotype observed in IGF-II null animals (Conover et al., 2004; De-Chiara et al., 1990). Although pappa is expressed in the mouse brain, a requirement for pappa in neural development, function, or behavior has not been reported (Conover et al., 2004). The zebrafish genome contains two pappa orthologs, pappaa and pappab, and both have been shown to cleave IGFBP-4 (Kjaer-Sorensen et al., 2013). Furthermore, morpholino knockdown of pappab affects developmental growth independent of its proteolytic activity (Kjaer-Sorensen et al., 2013). In contrast, the role of zebrafish pappaa has not been examined.

We find that the startle habituation phenotype in *pappaa/unfiltered*^{p170} mutants is caused by a mutation in *pappaa* and that *pappaa* promotes startle habituation through its metalloprotease activity (Figures 1B and 5). Our results support a mechanism in which PAPP-AA regulates habituation by targeting IGFBPs for cleavage and thereby enhancing IGF1R signaling. IGF1R signaling serves extensive roles during all stages of circuit development and also regulates circuit plasticity (Fernandez and Torres-Alemán, 2012). Although we cannot exclude the possibility that *pappaa* might contribute to some aspects of neural development, the ability to restore habituation in *pappaa* p^{170} mutants by acute activation of Akt and PI3K combined with the observation that acute inhibition of IGF1R reduces startle habituation strongly argues for an acute, post-developmental role for PAPP-AA during habituation learning.

Post-developmental roles for IGF1-IGF1R in learning behavior have been documented from C. elegans to rodent models of IGF signaling deficiency (Castro-Alamancos and Torres-Aleman, 1994; Tomioka et al., 2006). For example, insulin receptor signaling in C. elegans has been shown to play a critical role in experience-dependent temperature habituation and in taste avoidance learning (Ohno et al., 2014, 2014). Furthermore, in rats administration of insulin-like growth factor II enhances memory retention and prevents forgetting (Chen et al., 2011), and in humans insulin treatment has been reported to improve memory function (Benedict et al., 2004), while IGF-1-mediated signaling is thought to contribute to age-related cognitive decline (Deak and Sonntag, 2012). Reduced IGF availability and signaling has also been linked to schizophrenia (Venkatasubramanian et al., 2007), a disorder with a pronounced habituation deficit (Braff et al., 1992). This association warrants further examination of PAPP-A as a potential therapeutic target to stimulate IGF signaling in patients suffering from disorders like schizophrenia.

pappaa is expressed by neurons known to modulate startle behaviors and as outlined above is likely to promote habituation learning through an acute, IGFR-dependent process. IGFR signaling is known to regulate synaptic strength through ion channel modulation and neurotransmitter release, trafficking, and receptor activity (Blair and Marshall, 1997; Chen and Roche, 2009; Liou et al., 2003; Wang and Linden, 2000; Xing et al., 2007). In the future, resolving the precise mechanism by which *pappaa* modulates IGFR signaling and how IGFR signaling in turn promotes plasticity will be critical to understand the process of habituation leaning.

EXPERIMENTAL PROCEDURES

Fish Maintenance and Mutagenesis

ENU mutagenesis was performed in TLF and WIK *Danio rerio* strains as previously described (Dosch et al., 2004; Mullins et al., 1994). Embryos/larvae were maintained on a 14/10 hr light/dark cycle at 29°C and raised as previously described (Gyda et al., 2012; Kimmel et al., 1995). Behavioral experiments were conducted on 5–12 dpf larvae.

Behavioral Assays and Behavioral Analysis

Behavioral experiments were performed and analyzed with the FLOTE software package as previously described (Burgess and Granato, 2007a, 2007b; Hao et al., 2013; Wolman et al., 2011). The acoustic startle habituation assay was performed as described in Figure 1A (Wolman et al., 2011). Short latency C-bend startle response is measured by defined kinematic parameters including C-turn initiation latency, C-turning angle, C-turn duration, and Cturn maximum angular velocity (Burgess and Granato, 2007b). For secondary behavioral analyses of startle sensitivity, baseline activity, and visual habituation, habituation mutant larvae and wild-type siblings were identified based on acoustic startle habituation percentage (mutants < 50%, siblings > 50% habituation). Startle sensitivity assay was built into the habituation assay (Wolman et al., 2011). Baseline activity was determined by measuring total distance traveled over 160 s. For visual habituation testing, larvae were grouped by startle habituation phenotype in a 6 cm petri dish at a density of 12–15 larvae per dish. The visual habituation assay consisted of exposing larvae to 10 1-s-long dark flashes at 30 s ISI to establish baseline dark-flash responsiveness and then 30 1-s-long dark flashes at 3 s ISI to test visual habituation.

Recombination Mapping, Whole-Genome Sequencing, and Molecular Cloning of *pappaa* and *pcxa*

A three-generation mapping cross was built into our breeding scheme by crossing F1 heterozygotes from mutagenized TLF and WIK backgrounds. Pools of 25 behaviorally identified F3 mutant and sibling larvae were collected at 5 dpf and used for bulk segregant mapping with simple sequence length polymorphic markers (Table S1) (Jain et al., 2011) and/or whole-genome sequencing.

For whole-genome sequencing, we used 100 base pair paired-end sequencing on the Illumina HiSeq 2000 platform and compared gDNA from our pooled F3 mutants to gDNA prepped from the ENU-mutagenized males of both the TLF and WIK-L11 strains. Each sample was given its own lane in the flow cell, allowing us to achieve an average ${\sim}20\times$ coverage of the genome for each sample. Sequence data for each sample was mapped to the zebrafish Zv9 assembly (Ensembl) using Burrows-Wheeler Aligner software to map reads (Li and Durbin, 2009). Duplicate reads were eliminated with Picard tools (http://picard.sourceforge.net). We then adapted a fast homozygosity mapping strategy from Voz et al. (2012) to our screen. First we isolated all of the 100% homozygous bases unique to our TLF and WIK-L11 reference using the GATK software (McKenna et al., 2010). This generated 463,379 SNP markers. Using this set of SNPs, we calculated homozygosity scores by assigning TLF alleles a value of 1 and WIK alleles a value of 0 in a rolling 100-SNP window incrementing one SNP at a time. We analyzed the homozyogosity of our mutant samples across the genome and identified regions with scores >0.9 or <0.1 to indicate strong linkage to TLF or WIK-L11 alleles, respectively. To identify potentially causative mutations, we isolated all SNPs in the linked region that were unique to the mutant sample by comparing each mutant sequence to a combined reference sequence comprised of our TLF and WIK sequences, the Ensembl reference sequence, and our other mutant seguences. We restricted our list of candidate mutations to SNPs with <1% allele frequency in this reference sequence that also causes a change in amino acid sequence (nonsense, missense, or splice site mutations). For both unfiltered^{p170} and information overload^{p171}, these criteria produced a single SNP candidate.

To confirm candidate nonsense mutations in *pappaa* and *pcxa*, cDNA was prepared from total mRNA extraction from 5 dpf larvae, as previously described (Peterson and Freeman, 2009). *pappaa* and *pcxa* cDNA were amplified with primers (Table S1) designed against *pappaa* and *pcxa* reference sequence (Ensembl) with the following RT-PCR conditions: 94°C for 3 min and then 35 cycles of 94°C for 30 s, 57°C for 1 min, and 70°C for 1 min. Products were gel purified and cloned into the pCR2.1-TOPO-TA vector for sequencing.

For zebrafish *pappaa* RNA injection, cDNA was prepared from wild-type TLF larvae and amplified with the *z-pappaa:FL* primers using similar PCR conditions to those above but with extension time increased to 3 min. Full-length *pappaa* was cloned into pCS2⁺ vector, transcribed using the mMessage mMachine kit (Ambion), and injected at the one-cell stage at doses ranging from 1–200 picograms. Full-length human *pappa* and *pappa^{E483A}* constructs (Boldt et al., 2001) were similarly prepped and injected. Embryos injected with greater than 50 pg of zebrafish *pappaa* or 100 pg *h-pappa or h-pappaa^{E483A} mRNA* showed gross morphological abnormalities and necrosis, whereas embryos injected with 50 pg *pappaa* or 100 pg *h-pappa* (or less) appeared morphologically normal.

Genotyping, confocal imaging, morphological analyses, immunolabeling and in situ hybridization, pharmacological applications, and statistical analyses are described in Supplemental Experimental Procedures.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, five figures, one table, and three movies and can be found with this article online at http://dx.doi.org/10.1016/j.neuron.2015.02.025.

AUTHOR CONTRIBUTIONS

M.A.W., R.A.J., K.C.M., J.B.H., and M.G. designed research; M.A.W., R.A.J., K.C.M., H.B., J.S., and K.E.H performed research; M.A.W., R.A.J., K.C.M., K.E.H., J.B.H. and M.G. analyzed data; M.A.W. and M.G. wrote the paper.

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