Mutation of zebrafish dihydrolipoamide branched-chain transacylase E2 results in motor dysfunction and models maple syrup urine disease

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SUMMARY
Analysis of zebrafish mutants that demonstrate abnormal locomotive behavior can elucidate the molecular requirements for neural network function and provide new models of human disease. Here, we show that zebrafish quetschkommode (que) mutant larvae exhibit a progressive locomotor defect that culminates in unusual nose-to-tail compressions and an inability to swim. Correspondingly, extracellular peripheral nerve recordings show that que mutants demonstrate abnormal locomotor output to the axial muscles used for swimming. Using positional cloning and candidate gene analysis, we reveal that a point mutation disrupts the gene encoding dihydrolipoamide branched-chain transacylase E2 (Dbt), a component of a mitochondrial enzyme complex, to generate the que phenotype. In humans, mutation of the DBT gene causes maple syrup urine disease (MSUD), a disorder of branched-chain amino acid metabolism that can result in mental retardation, severe dystonia, profound neurological damage and death. que mutants harbor abnormal amino acid levels, similar to MSUD patients and consistent with an error in branched-chain amino acid metabolism. que mutants also contain markedly reduced levels of the neurotransmitter glutamate within the brain and spinal cord, which probably contributes to their abnormal spinal cord locomotor output and aberrant motility behavior, a trait that probably represents severe dystonia in larval zebrafish. Taken together, these data illustrate how defects in branched-chain amino acid metabolism can disrupt nervous system development and/or function, and establish zebrafish que mutants as a model to better understand MSUD.

INTRODUCTION
Maple syrup urine disease (MSUD) is an inherited metabolic disorder of branched-chain amino acids (BCAA): isoleucine, leucine and valine (Strauss and Morton, 2003; Chuang et al., 2006; Chuang et al., 2008). It demonstrates an autosomal recessive pattern of inheritance, and affects ~1 in every 185,000 children worldwide. However, much higher incidence rates are observed in Old Order Mennonite communities, with a ratio of 1:200 children, owing to founder effects. The first step of BCAA metabolism consists of a reversible transamination reaction to yield α-keto acids. The second step is oxidative decarboxylation of the α-keto acids by the mitochondrial branched-chain α-keto acid dehydrogenase (BCKD) complex. Mutation in any of the four genes encoding the three catalytic components of the BCKD complex has been shown to cause MSUD. Affected individuals accumulate BCAAs and α-keto acids in tissues and plasma, which cause the urine and bodily secretions to smell like maple syrup (burned sugar), hence the name of the disorder. In the most severe or ‘classic’ form of MSUD, the elevated BCAAs and α-keto acids can have a devastating impact on the central nervous system (CNS). If not treated, toxic build up of BCAAs and α-keto acids cause severe dystonia, coma, cerebral edema, dysmyelination and death within a couple weeks after birth. Treatment for MSUD typically consists of severe dietary restriction of BCAAs (Strauss et al., 2010). However, even with a carefully monitored diet, secondary illnesses can lead to metabolic crisis and neurological damage, which is thought to generate the mental retardation and psychiatric problems that are often observed. An alternative treatment for MSUD is elective liver transplantation, which reduces plasma levels of BCAAs and α-keto acids and spares the CNS from severe injury (Strauss et al., 2006). Unfortunately, the scarcity of livers for transplantation, surgical risks and lifetime usage of immunosuppressants limit this treatment option. A better understanding of the mechanisms that cause the neuropathology of MSUD and development of new therapeutic drugs could greatly benefit affected individuals as well as yield new insight into CNS metabolism.

To investigate the pathophysiology of MSUD, a mouse model of the classic form of the disease was created by deleting the gene encoding the E2 subunit of the BCKD complex (Homanics et al., 2006). A model of a less severe form of the disease, intermediate MSUD, was also developed. These mice recapitulate several aspects of MSUD, including elevated levels of BCAAs and α-keto acids in tissues and plasma, severe neurological impairment, and decreased phenotypic severity in response to liver cell transplantation (Skvorak et al., 2009a; Skvorak et al., 2009b; Zinnanti et al., 2009). However, the neuropathological mechanisms of MSUD remain poorly understood. Additional animal models could provide a complementary tool to study the mechanisms of this disease that generate CNS injury.
Developing zebrafish provide an excellent system to investigate the cellular and molecular mechanisms of MSUD. Although BCAA metabolism has yet to be described in this system, larval zebrafish offer several advantageous features, including small size, rapid development external to the mother, optical transparency, large clutch sizes and organ systems that are broadly similar to those in mammals. These features make zebrafish larvae readily amenable to a wide variety of behavioral, genetic, imaging, physiological and pharmacological approaches. The identification of zebrafish mutants with impaired BCKD complex function hold promise as a particularly useful tool to investigate the effects of disrupted BCAA metabolism.

Zebrafish perform a characteristic sequence of motor behaviors during development, which reflects different stages of locomotor network formation (Kimmel et al., 1974; Saint-Amant and Drapeau, 1998; Downes and Granato, 2006; McKeown et al., 2009). These behaviors were used as the basis for extensive mutagenesis screens to identify mutants with specific defects in embryonic motility (Granato et al., 1996). Mutants that demonstrated similar, abnormal behavior were grouped into phenotypic classes, including the accordion class. Starting around 21 hours post-fertilization (hpf), wild-type embryos demonstrate smooth, alternating tail coils. By contrast, accordion class mutants compress along the rostro-caudal axis and relax, like an accordion. Quietschkommode (que) mutants were grouped into the accordion class but, outside of the initial study identifying these mutants, it has not been characterized nor has the identity of the mutated gene been reported.

In this study, we reveal that que mutants contain a mutation that disrupts the E2 subunit of the BCKD complex. Correspondingly, we present evidence that que mutants accumulate high concentrations of BCAAs. que larvae also contain reduced levels of the neurotransmitter glutamate, which probably contributes to the aberrant CNS function and abnormal behavior observed in this mutant. Combined, these data identify the que mutant as a vertebrate model of MSUD, which holds promise as a powerful tool to better understand this disease and identify therapeutic compounds.

RESULTS
que mutants exhibit abnormal rostro-caudal compressions in response to touch
A single allele of que (ti274) was identified from a previously performed mutagenesis screen (Granato et al., 1996; Zottoli and Faber, 2000). The mutation is recessive, and homozygous mutants first demonstrate abnormal rostro-caudal compressions at around 72 hpf (data not shown). The abnormal behavior becomes more robust by 96 hpf. At this time point, high-speed video analysis shows that wild-type larvae respond to touch by first performing a large-amplitude body bend in which the nose touches the tip of the tail, a so-called C-start or C-bend (defined here as greater than 110°), followed by lower-amplitude body undulations to swim away (Fig. 1A; supplementary material Movie 1) (Eaton et al., 1977; Zottoli and Faber, 2000). By contrast, que mutants do not perform the initial C-bend or lower-amplitude body undulations, and instead demonstrate rostro-caudal shortening (Fig. 1B; supplementary material Movie 2). This behavior, called accordion behavior (Granato et al., 1996), is due to abnormal coordination of axial left-right muscle contractions. Other mutants have been shown to demonstrate accordion behavior due to defects in either CNS function, such as bandoneon (beo), or muscle relaxation, such as accordion (acc) (Gleason et al., 2004; Hirata et al., 2004; Hirata et al., 2005; Olson et al., 2010). We used kinematic analysis to compare the swimming behavior of wild-type, beo, acc and que larvae to each other to determine whether we could distinguish defects in CNS function from defects in muscle relaxation and classify que mutants. No clear trend emerged; however, que mutants consistently show the most dramatic disruption of swimming behavior compared with acc or beo mutants. Although acc and beo mutants demonstrate abnormal swimming behavior, C-bends are often observed (Fig. 1C-E). By contrast, que mutants very rarely execute large-amplitude body bends (Fig. 1F). que mutants continue to perform accordion behavior 5-6 days post-fertilization (dpf). They fail to inflate the swim bladder, which would enable them to feed, and eventually die around 7 dpf.

que mutants demonstrate abnormal motor output
To better examine whether que mutants harbor a defect in CNS function as opposed to a defect in muscle relaxation, we analyzed fictive locomotor output from the spinal cord by performing extracellular peripheral nerve recordings. Zebrafish larvae demonstrate bouts of motor output in response to touch (Fig. 2A,B). These bouts are composed of tightly coordinated bursts that alternate rapidly between the left and right sides and orchestrate the axial muscle contractions that constitute swimming. In wild-type larvae, we observed rapid alternations in locomotor output between the left and right sides, with little overlap in bursting activity (6.5±6%, n=5; Fig. 2C), as described previously (Masino and Fetcho, 2005). In que mutants, although the coordination of left-right locomotor activity was similar to wild-type siblings (compare Fig. 2C with 2D), the amount of overlap between left-right bursting activity was significantly increased (19.1±11.3%, n=7, t=-2.3, P<0.05). This increase in activity overlap is consistent with the abnormal coordination of left-right muscle contractions performed by que mutants. These data do not rule out the possibility that que mutants contain a defect in muscle relaxation; however, they do indicate that abnormal motor output from the CNS at least contributes to the behavior of this mutant.

To further characterize potential differences in locomotor output between wild-type and que mutant siblings, we examined a range of bout and burst properties related to rhythmic locomotor activity during fictive swimming (Masino and Fetcho, 2005). Although most of these properties were not significantly different between wild-type and mutant larvae (Table 1), que mutants generated a significantly greater number of bouts following touch stimulus applied to the head than did wild-type larvae [11.6±6.2 bouts and 1.9±0.9 bouts, respectively (t=3.5, P<0.01, n=5)]. These results suggest that, compared with wild type, there are subtle, yet significant, changes in the locomotor circuit that underlie swimming in que mutant larvae and participate in generating the mutant behavioral phenotype.

The que gene encodes dihydrolipoamide branched-chain transacylase E2 (Dbt)
To determine the molecular identity of the que gene, we used a positional cloning strategy. Using a three-generation map cross panel, we screened pools of genomic DNA from wild-type siblings.
and homozygous mutants with a panel of simple sequence length polymorphism (SSLP) markers. que mapped to chromosome 22, which confirmed previous low-resolution mapping results (Geisler et al., 2007). We then used DNA extracted from single embryos and single nucleotide polymorphism (SNP) markers to refine the map position to a 0.36 cM interval between the markers ENSDART109865 and wu:f63d09 (Fig. 3A). Extensive genome database analysis and sequencing of nearby candidate genes led us to dihydrolipoamide branched chain transacylase E2 (dbt), which encodes a subunit of the BCKD complex, which is required for BCAA metabolism. Zebrafish Dbt is a predicted 493 amino acids in length, and it is ~78.2% identical to the human protein (data not shown). Mutations in the human DBT gene are known to cause MSUD, which can result in severe dystonia and death if not treated. Given that que mutants demonstrate abnormal behavior and nervous system function consistent with the severe dystonia observed in humans, we sequenced the dbt gene. Sequence analysis of the dbt gene from que homozygotes revealed a single nucleotide substitution in the splice donor site of exon 6 compared with wild type. The guanine of the intron side of this splice site is changed to an adenine (Fig. 3B). To determine whether this change affects the splicing of intron 6, as would be predicted, we performed reverse

Table 1. Comparison of bout and burst properties related to fictive locomotor activity in wild type and que mutants

<table>
<thead>
<tr>
<th></th>
<th>Wild type</th>
<th>que mutants</th>
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<tbody>
<tr>
<td>No. of bouts</td>
<td>1.9±0.9</td>
<td>8.6±7.3*</td>
</tr>
<tr>
<td>Bout duration (msec)</td>
<td>329.1±60.4</td>
<td>285.2±148.8</td>
</tr>
<tr>
<td>No. of bursts per bout</td>
<td>9.5±1.9</td>
<td>9.7±3.6</td>
</tr>
<tr>
<td>Burst duration (msec)</td>
<td>13.3±2.8</td>
<td>12.4±1.7</td>
</tr>
<tr>
<td>Burst duty cycle (%)</td>
<td>30.9±11.8</td>
<td>39.6±6.8</td>
</tr>
<tr>
<td>Burst frequency (Hz)</td>
<td>31.6±4.6</td>
<td>37.6±9.3</td>
</tr>
<tr>
<td>Mean contralateral phase (%)</td>
<td>49.6±1.2</td>
<td>50.4±7.7</td>
</tr>
<tr>
<td>Bursting overlap (%)</td>
<td>6.5±6.0</td>
<td>19.1±11.3**</td>
</tr>
</tbody>
</table>

All values presented as mean ± s.d. Significant differences are indicated by *P<0.05 or **P<0.01.
contains an acetyl transferase domain, essential for its function that diminishes or abolishes BCKD complex function. (Lys257) was reported in an individual with the most severe or prematurely truncates the DBT protein at the orthologous position mutant protein. Interestingly, in humans, a mutation that prematurely truncate the Dbt protein by 224 amino acids. Downstream of Lys268 and contains four stop codons, which would downregulated transcripts (RT)-PCR using one primer in exon 6 and one primer in exon 7 (Fig. 3C). We found that RNA extracts from wild-type larvae were spliced according to prediction; however, RNA extracts from homozygous mutants revealed a larger transcript that contained the entire 86 base pairs of intron 6, indicating that it was spliced incorrectly (data not shown). This intron alters the sequence downstream of Lys268 and contains four stop codons, which would prematurely truncate the Dbt protein by 224 amino acids. 

To further confirm the molecular identity of que, we injected wild-type embryos with either a standard control morpholino or a morpholino designed to block translation of dbt. Embryos were injected at the one- to four-cell stage and monitored over the course of development. Embryos injected with the standard control morpholino exhibited mostly normal behavior throughout the course of development, is reminiscent of another mitochondrial protein progressive enrichment in the brain and gut organs over the course of development. Embryos were injected with the standard control morpholino designed to target dbt demonstrated clear rostro-caudal compressions and fewer large-amplitude body bends at 96 hpf, similar to que mutants (Fig. 3E). It is important to note that morpholinos are known to lose effectiveness at ~4-5 dpf owing to turnover, which probably explains why not all embryos injected with the dbt morpholino demonstrated the robust accordion behavior performed by que mutants (Bill et al., 2009). We also attempted to perform rescue experiments in mutant embryos by injecting mRNA encoding dbt at the one- to four-cell stage and analyzing motility behavior at 4 dpf. We did not observe rescue (data not shown); however, mRNA is known to lose effectiveness ~2 dpf owing to turnover. The que behavioral phenotype is not apparent until 3-4 dpf, which indicates that Dbt is required at this stage of development and precludes mRNA rescue. Regardless, the mapping data, nature of the que mutation, aberrant mRNA splicing observed in mutants, and morpholino phenocopy all argue that the que gene encodes Dbt.

dbt mRNA becomes enriched in the brain and gut organs during development

We next examined the spatial and temporal expression of dbt in developing zebrafish. RT-PCR revealed that dbt mRNA was present at all time points examined from 6 hpf to 120 hpf (Fig. 4A). In situ hybridization also confirmed early expression. dbt was detected at the two-cell stage, indicating it is a maternally deposited mRNA (Fig. 4B). The spatial expression of dbt is initially widespread through 24 hpf (Fig. 4C,D); however, its expression pattern over the next few days of development becomes enriched in the brain and organs in the gut, such as liver and intestine (Fig. 4E-H). These data suggest that dbt plays an important role in BCAA metabolism through function in these tissues. The prominent expression within the brain, in particular, suggests that dbt is important for CNS function. Intriguingly, the expression pattern of dbt, with progressive enrichment in the brain and gut organs over the course of development, is reminiscent of another mitochondrial protein that is important for CNS function, Opa3 (Pei et al., 2010).

que mutants harbor elevated levels of BCAAs

In mammalian systems, impaired DBT function, as demonstrated by MSUD-affected individuals, results in elevated levels of BCAAs (Strauss and Morton, 2003; Chuang et al., 2006; Chuang et al., 2008). Because dbt is disrupted in que mutants, we investigated their free amino acid profiles. Owing to the small size of larval zebrafish, a homogenate of 50 whole animals was used for each assay. We compared the free amino acid levels of wild-type and que mutant larvae at 96 hpf. Strikingly, que larvae harbor elevated levels of BCAAs (Fig. 5A,B). Isoleucine, leucine and valine concentrations were 788%, 1006% and 688% (n=3, P<0.01) of those of wild type, respectively. que mutants also showed a marked decrease in free glutamine levels, at 24% of wild-type (n=3, P<0.01). In addition,
Maple syrup urine disease zebrafish model

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Statistically significant decreases were observed in the levels of a wide variety of free amino acids, including aspartate (16%), GABA (32%) and serine (28% of wild type; all $n=3$, $P<0.01$); and alanine (44%), glutamate (38%), glycine (41%), methionine (49%) and threonine (60% of wild type; all $n=3$, $P<0.05$). To rule out the possibility that abnormal motor behavior itself alters free amino acid levels, we examined beo mutants, which contain a mutation in the glycine receptor $\beta_{2}$ subunit and exhibit abnormal behavior similar to que mutants (Fig. 5C) (Hirata et al., 2005). The free amino acid levels in beo mutants ($n=1$) were similar to that of wild-type controls, indicating that accordion behavior alone does not substantially alter free amino acid concentrations. Combined, these data provide strong evidence that mutation of the que gene results in an error in amino acid metabolism, yielding a prominent accumulation of BCAAs.

Glutamate levels are reduced in the brain of que mutant larvae

Although the neuropathology of MSUD is not well understood, reduced concentrations of neurotransmitters, including glutamate, were observed in the intermediate MSUD mouse model (Zinnanti et al., 2009). Neurotransmitter depletion was found to correlate with abnormal motor behavior and a highly abnormal posture consisting of recumbency and stiff, extended limbs. Given that our analysis of the free amino acids levels in que mutants showed decreased concentrations of free glutamate and these mutants demonstrate abnormal CNS function and motor behavior, we examined the distribution of glutamate using an antibody. As a control for antibody penetration and overall tissue morphology, we also stained using an acetylated tubulin antibody. Antibody penetration and general morphology of the brain of que mutants seemed similar to wild type at 96 hpf (compare Fig. 6A with 6D). By contrast, glutamate levels were markedly reduced in que mutant larvae ($n=5$ embryos, 12 sections, $P<0.01$; compare Fig. 6B with 6E, Fig. 6G), which probably contributes to the abnormal nervous system function and behavior observed by this stage of development.

DISCUSSION

In this study, we revealed that the dbt gene plays an essential role in developing zebrafish. The molecular nature of the que mutation, phenocopy through antisense morpholino injection, and the profile of free amino acid concentrations indicate that loss of dbt function result in abnormal amino acid metabolism and a dramatic accumulation of BCAAs. We determined that the dbt gene becomes enriched in the brain and organs in the gut during zebrafish development.
Disease Models & Mechanisms

Human systems, suggest a model for how mutation of the zebrafish
Development, and that impaired dbt function results in reduced
levels of glutamate in the brain. Because glutamate is crucial for
CNS function, the reduced levels of this neurotransmitter probably
promote the abnormal spinal cord output and accordion behavior
demonstrated by que mutants.

**dbt is required for brain function in zebrafish**

The findings from this study, as well as observations in rodent and
human systems, suggest a model for how mutation of the zebrafish
dbt gene leads to abnormal swimming. In mammalian systems, dbt
is required for the second step of BCAA metabolism, and its
impairment leads to elevated levels of BCAAs and \( \alpha \)-keto acids in
plasma and tissue (Chuang et al., 2006; Chuang et al., 2008). Elevated BCAAs in the plasma, in particular leucine, are thought
to out-compete other amino acids at the blood-brain barrier,
which results in neurotransmitter deficiencies, growth restrictions,
cytotoxic edema, myelin disruption, and impaired energy
metabolism throughout the CNS (Zinnanti et al., 2009).

Intracranial injection of \( \alpha \)-ketovaleric acid, which is derived from
valine, has been shown to elicit seizures in rats, whereas
administration of other \( \alpha \)-keto acids had no behavioral effect
(Choetinho et al., 2001). In individuals with MSUD, reducing the
concentrations of BCAAs and \( \alpha \)-keto acids in the plasma by liver
transplantation can protect CNS function and development
(Strauss et al., 2006).

We propose that very similar mechanisms regulate BCAA
metabolism in zebrafish (Fig. 7). Throughout the first 5 days
of zebrafish development, the embryo consumes the presumptive
equivalent of a high-protein diet in mammals by absorbing BCAA-
containing proteins from the yolk (Link et al., 2006; Tay et al., 2006).
During the earliest stages of development, within the first few hours
post-fertilization, the metabolic needs of the embryo are largely
met by maternally derived mitochondria and mRNA (Mendelsohn
and Gitlin, 2008; Zhang et al., 2008; Abrams and Mullins, 2009).

However, as embryogenesis proceeds, BCAA metabolism
increasingly relies upon zygotic transcription. In wild-type
embryos, BCKD complex function in the liver, other gut organs
and the brain itself, protect the CNS from BCAA toxicity, similar
to mammalian systems. According to this model, BCKD complex
function supports appropriate import of amino acids into the CNS,
robust metabolic generation of neurotransmitters, which are
essential to support the coordinated CNS output that generates
vigorous swimming behavior. In que mutants, our data indicate that
mutation of dbt disrupts BCKD complex function to cause the toxic
accumulation of BCAAs and, probably, \( \alpha \)-keto acids. This error
probably causes abnormal retention, metabolism and import of
amino acids into the CNS, reduced levels of glutamate and other
neurotransmitters, abnormal CNS function, and accordion
behavior (Fig. 7). It will be interesting to use transgenic approaches
to determine whether restoring gene function in the liver of que
mutants preserves normal CNS function, as shown in mammals
(Strauss et al., 2006; Skvorak et al., 2009a; Skvorak et al., 2009b).
Driving gene expression in other organs not yet explored in
mammals can also be investigated, which might indicate new
therapeutic options for individuals with MSUD.

**que mutants are a new animal model of MSUD**

*que* larvae harbor a mutation in zebrafish dbt, which results in
elevated BCAA levels, similar to both the mouse models of MSUD
and affected humans. In the mouse model of intermediate MSUD,
elevated levels of BCAAs have been shown to correlate with
progressive disruption of CNS function and concomitant defects
in motor behavior that culminate in severe dystonia (Silberman
et al., 1961; Morton et al., 2002; Zinnanti et al., 2009). Similarly,
severe dystonia has been reported in MSUD-affected individuals
during acute metabolic decompensation (Silberman et al., 1961;
Morton et al., 2002; Zinnanti et al., 2009). que mutants
demonstrate a progressive defect in motor behavior that culminates in abnormal CNS function and accordion behavior.
Accordion behavior is probably the expression of severe dystonia in developing zebrafish. We and others have previously shown that zebrafish mutants that exhibit accordion behavior contain mutations in genes known to control movement and muscle tone in mammalian systems (Downes and Granato, 2004; Gleason et al., 2004; Hirata et al., 2004; Hirata et al., 2005; Wang et al., 2008; Olson et al., 2010).

The findings from this study indicate that que mutants are a new animal model of MSUD. One aspect of MSUD is the distinct, maple syrup smell of bodily secretions of affected individuals. We did not detect any distinct odor of que mutants (data not shown); however, this is probably due to the small size and minute amounts of secretions produced by larval zebrafish. Nevertheless, que larvae seem to recapitulate molecular, biochemical, cellular and behavioral aspects of MSUD. Because larval zebrafish contain a smaller nervous system than do mammalian systems, with fewer numbers of cells, que mutants provide a promising system to better characterize the progression of CNS injury in response to BCAA toxicity. Moreover, the small size, aquatic nature, development that is external to the mother and the ability to obtain large numbers of zebrafish embryos make them amenable to small-molecule screens (Zon and Peterson, 2010). The behavioral phenotype of que mutants is robust and easily quantifiable; therefore, que mutants could be developed into a high-throughput system to screen libraries of compounds to identify small molecules that improve swimming behavior.
improve the behavioral phenotype of que mutants could be candidate therapeutics for individuals with MSUD.

Other aspects of MSUD can also be investigated using larval zebrafish. Gene targeting approaches can be readily employed, such as morpholino injection or zinc-finger nuclease technology, to model MSUD caused by disruption of other BCKD complex subunits (Ekker, 2008; Bill et al., 2009). These technologies can also be used to examine the in vivo role of BCKD regulatory proteins, such as the BCKD phosphatase or kinase. Genetic modifier screens can also be performed using the que mutant to search for genes such as the BCKD phosphatase or kinase. Genetic modifier screens can also be performed using the que mutant to search for genes that can compensate for disruptions in BCAA metabolism. Taken together, these approaches can provide a promising platform to better understand CNS metabolism and develop new therapies to combat MSUD.

METHODS
Zebrafish maintenance and breeding
All animal protocols were approved by the Institutional Animal Care and Use Committees (IACUC) at the University of Massachusetts and the University of Minnesota. Zebrafish were raised and maintained according to standard procedures. Developing zebrafish were kept at 28.5°C in E3 media and staged according to morphological criteria (Kimmel et al., 1995; Parichy et al., 2009). Experiments were performed using que, beo and acp mutant alleles maintained on a mixed Tübingen (Tü) or tub longfin (TLF) genetic background.

Behavioral analysis
To characterize swimming behavior, light-touch stimuli were applied to the head of larvae using a 1 mm insect pin. The response was recorded using a high-speed video camera (Fastec Imaging, San Diego, CA), recording 500-1000 frames per second, mounted to a 35 mm lens (Nikon, Melville, NY). The head-to-tail angle for each frame was measured using automated software developed by G.B.D’s laboratory (Kelly Anne McKeown and Sandy Whittlesey unpublished). Briefly, pixel density analysis was used to identify three landmarks along the larval body: the tip of the nose, the border between the yolk ball and yolk extension, and the tip of the tail. These three points form an angle, and these angles were plotted over time using Microsoft Excel.

Electrophysiological recordings
Zebrafish larvae at 4 dpf were anesthetized with 0.02% Tricaine-S (Western Chemical) in extracellular recording solution (Legendre and Korn, 1994; Drapeau et al., 1999; Masino and Fetcho, 2005) and paralyzed with 0.1% (w/v) α-bungatoxoin (Sigma), which significantly reduced or abolished postsynaptic muscle activity based on patch recordings from muscle fibers (Masino and Fetcho, 2005). The extracellular solution was superfused continuously at 22-26°C. Larvae were pinned in a dorsoventral position to a Sylgard-lined glass-bottom Petri dish and the skin was removed. Extracellular suction electrode recording techniques were used to monitor the activity of peripheral nerves during fictive behavior (Masino and Fetcho, 2005). Activity occurred spontaneously but was also initiated by gently applying a touch stimulus to the head with a tungsten pin controlled and positioned by a manual micromanipulator (MX130, Siskiyou, Grants Pass, OR). The tip of the extracellular suction electrode (~15 μm tip diameter) was positioned at the dorsoventral midline of a myotomal cleft where the skin had been removed. All extracellular recordings were restricted to between body segments 7 and 15. A MultiClamp 700B (Molecular Devices, Sunnyvale, CA) amplifier was used to monitor extracellular voltage in current-clamp mode at a gain of 1000 (Rf=50 MΩ) with the low- and high-frequency cut-off at 100 and 4000 Hz, respectively. Recordings were sampled at 10 kHz. Extracellular recordings were digitized using a digitizing board (DigiData series 1440A, Molecular Devices, Sunnyvale, CA) acquired using pClamp 10 software and rectified offline.

Analysis of peripheral nerve activity
A program written in MATLAB (Mathworks, Natick, MA) was used to analyze the data. Estimates of mean burst frequency were determined from a Fourier transform, such that the initial estimate of mean burst frequency was the frequency at which the Fourier transform magnitude peaked over a frequency band from 0.1 to 5 Hz. The rectified voltage recordings were smoothed with a Gaussian-weighted moving average with 99% of the weight...
concentrated over an interval whose width was one-quarter of the reciprocal of the estimated burst frequency (quarter-width'). The occurrence times of rhythmic bursts in the smoothed voltages were determined with an algorithm that searched for local peaks and troughs over quarter-width intervals while forcing adjacent peaks and troughs to be separated by at least quarter-width, and furthermore forcing peaks and troughs to alternate. With the peaks and troughs defined, the individual 'burst sections' were then defined as the interval between adjacent troughs. To determine the start of individual bursts, the burst-onset was defined as the time at which the smoothed waveform rose from the first trough to 10% of the way to the next peak. Similarly, burst-termination was defined as the time at which the smoothed waveform fell from the peak by 90% of the vertical distance to the next trough. Next, the analysis program was used to determine the bout and burst properties for each voltage trace in a manner similar to that used by Masino and Fetcho (Masino and Fetcho, 2005). Finally, the amount of overlap in activity between alternating (left-right) bursts was measured as the proportion of the total burst time occupied by the simultaneous activity in the paired (left-right) extracellular recordings (Fig. 2C,D, overlap indicated by gray bars). The means and standard deviations for each parameter were then determined.

Chromosomal mapping and sequence analysis
CROSSES BETWEEN FISH HETEROZYGOS FOR THE que Allele AND WIK Fish were used to generate a three-generation map cross panel. F2 que mutant embryos and wild-type siblings were collected, sorted based upon the 96 hpf phenotype and stored in methanol at −20°C. DNA was extracted from more than 833 mutant larvae, and SSLP markers and SNP markers were obtained and generated against genes to refine the mapping interval. Exons and intron-exon boundaries of candidate genes were sequenced (Genewiz, South Plainfield, NJ) from wild type, que siblings and homozygous mutants.

Morpholino analysis
WILD-TYPE ZEBRAFISH EMBRYOS were pressure injected at the one- to four-cell stage with 12 ng of morpholino designed to block translation of dbt or the standard control morpholo (Gene Tools, Phinomath, OR). This amount of morpholino was selected based upon dose- response experiments in which higher doses were found to generate morphological defects and/or lethality. The sequence of the translation-blocking morpholino was 5’-CGCAGTAAATGACCACCGCCCATCT-3’. Underlined residues indicate the start codon. The control morpholino sequence was 5’-CCTCTTACCTCAGTTACAATTATA-3’. The embryos were raised at 28.5°C, and locomotive behavior was examined across development. Kinematic analysis, as described above, was performed at 96 hpf.

RT-PCR
RT-PCR was used to analyze mRNA splicing in mutants as well as examine expression during development. Primers designed against dbt protein-coding exon 6 (5’-ATCAAACTAACGGAAGTTGTCCG-3’) and exon 7 (5’-CGCAGTCCGACCCACGCAA-3’) were used to amplify cDNA from wild-type and homozygous que mutant larvae. The primers used to amplify β-actin were 5’-CACACGGTGCGCCATCTATGA-3’ and 5’-AGGATCTTCTATCAGGATTAGTCTTGG-3’. The RNAs were reverse transcribed using the Omniscript kit (Qiagen, Venlo, The Netherlands). The dbt PCR products were sequenced for confirmation (Genewiz, South Plainfield, NJ). RT-PCR reactions were performed multiple times to decrease the likelihood of amplification artifacts.

Whole-mount in situ hybridization
Anti sense digoxigenin probes were generated against dbt using cDNA (Genbank ID BC099017) acquired from Open Biosystems (Huntsville, AL). Whole-mount, colorimetric in situ hybridization was performed using established protocols (Thisse and Thisse, 2008) and examined using a compound microscope (Zeiss, Thornwood, NY) attached to a digital camera (Zeiss, Thornwood, NY). Cross-sections were generated by hand sectioning in situ hybridization stained embryos with a razor blade attached to a surgical blade holder. To generate sagittal sections, in situ hybridization stained embryos were embedded in 1.5% Agar and 5% sucrose. Blocks were kept in 30% sucrose solution overnight. The next day blocks were cut into 20 mm sections using a cryostat (Leica, Buffalo Grove, IL).

Amino acid quantification
For each amino acid quantification experiment 50 96-hpf larvae were sorted based upon the locomotor phenotype, were flash frozen in liquid nitrogen and stored at −80°C. The samples were homogenized and precipitated with 0.1 M lithium citrate, 3.3% 5-sulphosalicylic acid. The samples were sonicated for 10 minutes, then centrifuged at 4600 g for 20 minutes. The supernatant was then applied to VivaSpin500 size exclusion columns (Sartorium, Germany) and centrifuged at 15,000 g for 4 hours. The flow-through was stored at −80°C then sent to the University of California Davis California Genome and Proteomics Center to resolve free amino acid concentrations.

Clinical issue
Maple syrup urine disease (MSUD) is an inherited disorder that results in disrupted metabolism of branched-chain amino acids (isoleucine, leucine and valine), resulting in the toxic accumulation of these amino acids and their by-products. This disease can have a devastating impact on the central nervous system (CNS), resulting in mental retardation, severe dystonia, coma or death if not treated. Although mouse models of the disease have been developed and some disease genes are known, the cellular and molecular mechanisms that promote brain injury in individuals with MSUD are not well understood.

Results
In this paper, the authors characterize a zebrafish mutant called quetschkommode (que) that exhibits defects in motor behavior. The mutation is found in the dihydrolipoamide branched chain transacylase E2 (dbt) gene, a homolog of the human DBT gene, which can cause MSUD when mutated. In addition to abnormal behavior, que fish are shown to have disrupted metabolism of branched-chain amino acids and aberrant CNS function, mirroring features of human MSUD.

Implications and future directions
These data reveal the que zebrafish mutant to be a new animal model of MSUD. Because zebrafish offer a number of advantages for cellular, molecular, pharmacological and genetic analysis, the que mutant provides a unique tool to deepen our understanding of and develop new therapeutic options for this disease.
Antibody staining
The tissue was prepared by embedding 96 hpf larvae as described for in situ hybridization. Double immunostaining was performed using standard protocols (similar to Downes and Granato) (Downes and Granato, 2004). The primary antibodies were rabbit anti-L-glutamate (1:100 dilution, ab9440; Abcam, Cambridge, MA) and mouse anti-acetylated tubulin (1:200 dilution, T6793; Sigma, St Louis, MO). Secondary antibodies were Alexa Fluor 488 goat anti rabbit (1:1000 dilution, A11034; Molecular Probes/Sigma, St Louis, MO) and Alexa Fluor 594 goat anti mouse (1:1000 dilution, A11005; Molecular Probes, Sigma, St Louis, MO). Images were acquired using a confocal microscope (Nikon, Melville, NY).

The fluorescent intensity for acetylated tubulin and L-glutamate antibody staining was quantified using the EZ Viewer program (Nikon, Melville, NY) by collecting entire frames (10,1283 μm2) or selecting a region of interest above the notochord (3060 μm2) for both channels. The numbers used for quantification are the analog-to-digital converter (ADC) values of L-glutamate normalized to acetylated tubulin.

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

AUTHOR CONTRIBUTIONS

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SUPPLEMENTARY MATERIAL
Supplementary material for this article is available at http://dmm.biologists.org/lookup/suppl/doi:10.1242/dmm.008383/-/DC1

REFERENCES
Disease Models & Mechanisms  ●  DMM

**Maple syrup urine disease zebrafish model**


