LRT, a tendon-specific leucine-rich repeat protein, promotes muscle-tendon targeting through its interaction with Robo

Bess Wayburn and Talila Volk*

Correct muscle migration towards tendon cells, and the adhesion of these two cell types, form the basis for contractile tissue assembly in the Drosophila embryo. While molecules promoting the attraction of muscles towards tendon cells have been described, signals involved in the arrest of muscle migration following the arrival of myotubes at their corresponding tendon cells have yet to be elucidated. Here, we describe a novel tendon-specific transmembrane protein, which we named LRT due to the presence of a leucine-rich repeat domain (LRR) in its extracellular region. Our analysis suggests that LRT acts non-autonomously to better target the muscle and/or arrest its migration upon arrival at its corresponding tendon cell. Muscles in embryos lacking LRT exhibited continuous formation of membrane extensions despite arrival at their corresponding tendon cells, and a partial failure of muscles to target their correct tendon cells. In addition, overexpression of LRT in tendon cells often stalled muscles located close to the tendon cells. LRT formed a protein complex with Robo, and we detected a functional genetic interaction between Robo and LRT at the level of muscle migration behavior. Taken together, our data suggest a novel mechanism by which muscles are targeted towards tendon cells as a result of LRT-Robo interactions. This mechanism may apply to the Robo-dependent migration of a wide variety of cell types.

KEY WORDS: Drosophila, Leucine-rich repeat, Robo, Muscle, Tendon

INTRODUCTION

The correct migration and adhesion between cells of different origin is essential for tissue morphogenesis. The development of the Drosophila embryonic contractile tissue established by the migration and adhesion of muscles towards their corresponding tendon cells has served as a model through which to study various aspects of tissue assembly, including guided muscle migration and the formation of the myotendinous junction (Schnorrer and Dickson, 2004; Volk, 1999). Somatic myotubes produced at distinct embryonic sites migrate towards specific ectodermal cells, termed tendon cells, defined in Drosophila by the expression of the transcription factor Stripe (Volk and VijayRaghavan, 1994; Williams and Caveney, 1980). Stripe is an EGR-like transcription factor that defines tendon cell identity in the ectoderm (Frommer et al., 1996). Stripe is both necessary and sufficient to promote muscle migration towards tendon cells (Becker et al., 1997; Volk and VijayRaghavan, 1994; Vorbruggen and Jackle, 1997). In addition, Stripe is required for the maturation of tendon cells following their attachment to muscles (Volohonsky et al., 2007). Subsequently, myotubes form integrin-mediated hemi-adherens-type junctions with tendon cells at both their ends so that each myotube is stretched between two ectodermal tendon cells (Bokel and Brown, 2002). This organization enables the typical wave-like movement of the hatched crawling larvae.

Muscle migration towards tendon cells is a multistep process in which founder cells first move apart from each other; then, during and following muscle fusion, each myotube directs its leading edge towards a specific tendon cell in response to guidance cues provided by the tendon (Schnorrer and Dickson, 2004; Volk, 1999). Often, the migrating myotube first establishes contact with a tendon cell located at the posterior segmental border, and then extends its opposite end towards a tendon cell located at the anterior segment border. Following the arrival of the muscle’s leading edge to this tendon cell, its migration is arrested and the myotendinous junction is formed.

Tendon cells are crucial in providing guidance for muscle migration. In the absence of such signals (e.g. in stripe mutants), muscles lose their directional migration and fail to attach to tendon cells, leading to a complete disruption of the muscle pattern and failure of the larvae to move (Frommer et al., 1996). Positive guidance cues produced by tendon cells that act to attract muscles towards tendon cells were described. These include proteins directing axon migration, such as Slit and Syndecan (Kramer et al., 2001; Steigemann et al., 2004). Signals mediating the arrest of muscle migration once it has reached its targeted tendon cell have not yet been described.

Slit is secreted by tendon cells under the regulation of Stripe (Kramer et al., 2001; Volohonsky et al., 2007), whereas Roundabout (Robo) and Robo2 (Leak – FlyBase) are expressed in specific muscles (Kramer et al., 2001). Slit expression is not unique to the muscle/tendon junction, and this protein also mediates axonal pathfinding in the central nervous system (Dickson and Gilestro, 2006). However, unlike the Slit-Robo interactions in the nervous system, where Robo appears to mediate the repulsion or turning away of the axonal leading edge, muscles respond to Slit in two distinct fashions (Kramer et al., 2001). The ventral oblique muscles, which migrate ventrally, are first repulsed from the ventral midline in response to Slit, and in later developmental stages they are attracted to Slit provided by the segmental border tendon cells. Consequently, in slit or robo mutant embryos, the ventral oblique muscles cross the ventral midline, resulting in an aberrant pattern. Midline rescue of Slit in slit mutant embryos exhibits a muscle phenotype in which the ventral longitudinal muscles fail to attach to the tendon cells at the segmental border tendon cells. In addition, ectopic expression of Slit in the two rows of cells in the Engrailed domain leads to attraction of the ventral longitudinal muscles to this ectopic site (Kramer et al., 2001).
In an effort to identify additional tendon-specific signals essential for the guidance of migrating muscles towards tendon cells, we recently performed a microarray screen for Stripe downstream targets. One of the genes recovered was CG11136, a novel gene that encodes a type 1 transmembrane protein containing a leucine-rich repeat (LRR) domain in its extracellular region. LRR domains are formed by a string of consecutive leucine-rich repeats separated by variable inter-strand segments. This domain has been implicated in protein-protein interactions (Bella et al., 2008). Several transmembrane or secreted LRR proteins have been described in Drosophila, including Tartan, Capricious and others (Dolan et al., 2007; Kurusu et al., 2008). Mutants in these genes exhibit a wide range of phenotypes, depending on their tissue-specific distribution. However, they share a common function in cell-cell recognition events (Blair, 2001; Kohsaka and Nose, 2009; Krause et al., 2006; Milan et al., 2002; Milan et al., 2005; Milan et al., 2001; Sakurai et al., 2007). Two additional LRR proteins, Haf and CGI8561 (Convolute – FlyBase), expressed by muscle cells, affect motor axon recognition of specific muscles (Kurusu et al., 2008). The molecular basis for the activity of these proteins is yet to be elucidated.

Here, we show that CG11136 is a tendon-specific LRR transmembrane protein that is required for the establishment of the correct embryonic muscle pattern. In its absence, specific muscles do not extend properly to their attachment sites, and other muscles do not arrest their migration behavior. Due to its tendon-specific distribution we named the CG11136 protein ‘Leucine-rich tendon-specific protein’ (LRT). We show that LRT forms a protein complex with Robo, and that these proteins functionally interact, thereby affecting the extent of muscle extension towards tendon cells. Taken together, our data suggest a model in which the association of LRT with Robo receptors at the level of the muscle-tendon junction is essential for better targeting of muscles to their tendon cells as well as arrest of muscle migratory behavior.

MATERIALS AND METHODS

Fly stocks
The following fly stocks were used: yw, stripe-gal4 (G. Morata, Madrid, Spain), mef2gal4, actin-gal4, UAS-slit, Df(2R)BSC403 and Df(2R)Exel1764 (Bloomington Stock Center); UAS-lrt-gfp and UAS-lrt-ntd-gfp were created in the lab (injected in flies by Genetic Services); UAS-CG8561-GFP and 5053-gal4/TM3 (a gift from Frank Schnorrer, Max Planck Institute of Biochemistry, Martinsried, Germany); 44991-expressing RNAi against LRT (Vienna Drosophila Research Center); robo/cyo slt/cyo (B. Dickson, IMP, Vienna, Austria); UAS-Robo-RNAi; Robo2-RNAi; Robo3-RNAi (a gift from V. Rodrigues, NCBS, Bangalore, India).

The fly lines Df(2R)BSC403/cyo;UAS-lrt-gfp and Df(2R)BSC403/cyo;sr-gal4/TM6 were created to enable rescue of Df(2R)BSC403. In all cases, a Kr-GFP marked CyO balancer was used to identify the homozygous mutants.

For the analysis of the phenotype of Muscle 12 we used flies with the following genotype:Df(2R)BSC403/Cyo,KrGFP; 5053-gal4/UAS-CG8561-GFP/TM6. The GFP pattern of the 5053-gal4 is distinct from that of the Kr-GFP, enabling the identification of the homozygous mutants.

For the rescue of the flies above we crossed Df(2R)BSC403, UAS-LRT-GFP/Cyo,Kr-GFP; 5053-gal4/TM6 X Df(2R)BSC403/Cyo,Kr-GFP; sr-gal4/TM6. The GFP was detected in three distinct sites: in tendon cells, in muscle 12 and in Kruppel (Kr) pattern. The homozygous rescued mutants were identified by negative staining of the Kr pattern, and positive GFP staining of muscle 12 as well as tendon cells.

Constructs
The Gateway Cloning Technology (Invitrogen Life Technologies, USA) was used to produce the following constructs: full lrt was amplified for insertion into the Gateway pDonr-201 vector with primers 5'-GGGGACGTTTTGACAAAAAAGCAGCTCTCGACGCTGATAAAGAGG-3' (forward) and 5'-GGGGACCATCTTGTGACAAAGAAGCTGGTTACGCTGCTGCTCGT-3' (reverse). The insert was then transferred to destination vector pTWG, which contains a UAS promoter sequence and a 3' GFP tag. This construct was used to drive GFP-labeled LRT expression in cells as well as in transgenic flies.

UAS-lrt-ntd-gfp was created with the same forward primer as the complete LRT, and the following reverse primer: 5’-GGGGACC-ACTTTGTACAAGAAGCTGGTTACGCTGCTGCTGCTCGT-3’. pUAST-Robo-HA construct was obtained from B. Dickson (IMP, Vienna).

5’ RACE of CG11136
The 5’-ends of mRNAs extracted from an embryo collection of 10- to 16-hour-old embryos were specifically capped with a known sequence. Primers specific to the cap as well as to the known region of LRT (at its 5’ region) were then used to amplify the potential unknown 5’ region. A fragment of 520 bp was identified that contained the 5’ UTR region of LRT as well as part of its coding sequence. This fragment was fused to the rest of LRT cDNA and was used in all further experiments (rescue, overexpression and immunoprecipitation).

In situ hybridization
A DNA probe was created with following primers: 5’-CCAAATGACCTCAAGGATCC-3’ (forward) and 5’-GTGTTGAGCCAGATCGCAT-3’ (reverse). In situ hybridization was performed as described (Subramanian et al., 2007).

Immunohistochemical reagents
The following primary and secondary antibodies were used: mouse anti-GFP antibody (Roche Diagnostics); mouse monoclonal anti-HA antibody (11583816001, Roche, Switzerland) at 1:200 for immunostaining and 1:1000 for western blot; anti-Myxosin heavy chain (P. Fisher, Stony Brook, NY, USA) at 1:500 for immunostaining. Anti βPS-integrin (Hybrida Center) was used at 1:2 for immunostaining. Guinea pig anti-Sr (produced in our lab) was used 1:200 for immunostaining. Mouse anti-Robo [Hybrida Center and S. Kramer (UMDNJ-Robert Wood Johnson Medical School, NJ, USA)] was used at 1:10 for immunostaining. Chick anti-GFP was used at 1:500 for immunostaining. Secondary antibodies conjugated with Cy3, Cy5 and FITC raised against guinea pig, rat, mouse and rabbit were purchased from Jackson ImmunoResearch Laboratories.

Rat anti-LRT (see Results) was raised against a unique sequence in the N-terminal region of the protein and was used at 1:200 for immunostaining.

SRR and S2 Schneider cell transfections
Drosophila S2 cells were maintained in Schneider’s Drosophila Medium (01-150-1A, Biological Industries, Israel) + 10% FBS (12657-029, Gibco, USA) at 25°C. Cells were transfected using the Electra IV transfection reagent (L3287, Sigma-Aldrich, USA) according to the manufacturer’s instructions. The transfected cells were incubated for 24 hours before use for protein extraction or immunostaining.

Immunofluorescent staining procedures
The embryos were dechorionated and fixed in 4% formaldehyde (FA). Following fixation, the embryos were devitellinated and blocked in 10% BSA in PTW for 1 hour. Embryos were incubated with the primary antibody over night. Then the embryos were washed with PTW, secondary antibody was added for 2 hours, and then washed again and dehydrated in 80% glycerol.

For heat fixation, the embryos were collected from the sieve using a spatula and added to 5 ml of a boiling solution of 0.7% NaCl and 0.04% Triton-X in a 50 ml Eppendorf tube. After 5 seconds, the tube was transferred to ice and 20 ml of ice-cold NaCl/Triton solution was added. The embryos were incubated on ice for 15 minutes and then transferred to a 2 ml Eppendorf tube. The Triton/salt solution was removed, and the vitelline membrane removed by vigorous shaking in a 1:1 heptane/methanol solution. The embryos were then incubated in methanol for at least 1 hour before normal dehydration and staining as described above.
Immunoprecipitation and western analysis
Transfected S2 cells were harvested by centrifugation in a 15 ml tube at 1200 RPM to remove the media. The pelleted cells were then washed with PBS. The cells were lysed in NP40 buffer (10 mM Tris-Cl pH 8.0, 150 mM NaCl, 0.1% NP40) + protease inhibitors (Sigma P 8340) and incubated on ice for 10 minutes. After pelleting the debris, the crude extract was added to Protein G PLUS-Agarose beads (sc-2003, Santa-Cruz Biotechnology, USA) preloaded with the anti-GFP antibody overnight at 4°C. The mixture was allowed to incubate overnight at 4°C in a rotating arm. The beads were pelleted, washed in NP40 buffer and boiled in sample buffer for use in the western analysis. Western analysis was performed according to standard procedures described (Volk, 1992).

Stripe microarray
Embryos aged 10-15 hours after egg laying were collected, either of wild type or carrying 69B-gal4-UAS-stripeB constructs. The total RNA was extracted using the Macherey-Nagel NucleoSpin RNA II mini kit, following the protocol, and then kept at –70°C. Total RNA was prepared independently five times from embryos of each genetic background in order to better normalize the age of these embryo populations. The RNA samples were then collected and concentrated to give 1 mg of total RNA using the RNA cleanup RNeasy Mini Kit (Qiagen). The probe preparation, cDNA synthesis, cRNA reactions and hybridization with Affymetrix high-density oligonucleotide arrays for Drosophila melanogaster was carried out in the Weizmann Institute microarray unit. The microarray experiment was repeated four times. CG11136 (LRT) was consistently elevated in the order of 20.8. In a microarray screen designed to identify genes that are expressed regulated by Stripe microarray, the expression of CG11136 gene, labeled for LRT (red) and showing no staining. Scale bars: 20 μm in C,D,G.

Confocal microscopy
Visualization was performed with a Zeiss LSM 510Meta or a Zeiss LSM710 confocal system. The fluorescent digital images were processed using Adobe PhotoShop CS.

RESULTS

LRT is a tendon-specific protein that is positively regulated by Stripe
In a microarray screen designed to identify genes that are expressed downstream of Stripe (for details see Materials and methods), we identified CG11136. The CG11136 open reading frame sequence contains a signal peptide, a leucine-rich repeat domain, a transmembrane domain and a small cytoplasmic domain at its C-terminal end, suggesting that it is a type I transmembrane domain protein (Fig. 1A). In situ hybridization with an antisense RNA probe for CG11136 indicated an expression pattern similar to that of Stripe, suggesting a tendon-specific distribution (Fig. 1B). Thus, we named the CG11136 gene product Leucine-rich tendon-specific protein (LRT). The mRNA expression of LRT was detected from embryonic stage 12 and persisted to late stage 16. Overexpression of Stripe in the Engrailed domain (driven by en-gal4) led to expression of lrt in the en-grailed domain, which is wider by an additional row of cells relative to the Stripe-positive cell row in the segmental border (Fig. 1C,D), indicating that lrt expression can be promoted by Stripe. However, we found that the protein coded by lrt was still expressed in stripe mutant embryos (data not shown), suggesting an earlier transcripational regulation of this gene, similar to some of the stripe downstream genes described previously (Chanana et al., 2007; Subramanian et al., 2007).

An antibody was raised against the unique extracellular domain of LRT, using as an antigen a recombinant fragment that excluded the LRR domain, to prevent cross-reactivity with other LRR proteins. Staining of embryos with this antibody revealed a prominent expression pattern of LRT specifically in tendon cells (Fig. 1E,F,G). The intensity of staining varied between the different types of tendons, but it appeared that all tendons were positive for LRT staining. LRT protein staining was detected only at relatively late embryonic stages (stages 15-16 and beyond) following the establishment of muscle-tendon attachment (Fig. 1E; Fig. 2). Tendon cells of embryos at stages 13-14, which do not yet bind to muscles, did not exhibit positive staining for LRT protein (not shown), although these cells expressed lrt mRNA (see Fig. 1B). It is possible that the antibody was not sensitive enough to detect low protein levels of LRT in these cells. The pattern of LRT staining appeared either as an oval patch on the tendon membrane (e.g. on the ventral longitudinal muscles; see Fig. 1E, Fig. 2A), or as a line (e.g. Fig. 2G). Significantly, LRT accumulated only at the sites of muscle tendon attachment (Fig. 2C, arrowhead), and not on tendon cells that do not bind muscles (Fig. 2C, arrow). To verify the localization of LRT along the tendon cell membrane, we double-labeled embryos with anti-LRT and a tendon-specific membrane marker (CD8-GFP). Single confocal optical sections of the tendons of these embryos showed that LRT was completely absent from the apical surfaces of the tendon cell (not shown) and was detected only at the basal
Fig. 2. LRT accumulates at the surfaces of the tendon cell corresponding to muscle ends. (A-C) Whole-embryo staining of a wild-type embryo labeled for Myosin heavy chain (MHC; green, A,C), LRT (red, A-C) and Stripe (blue, B,C), showing the association of LRT with the muscle ends. (D-I) Single confocal optical sections of an embryo expressing tendon-specific membrane CD8-GFP and stained for GFP (green, E,F,H,I), LRT (red D-I) and MHC (blue, F,I), showing a partial overlap between GFP and LRT. LRT distribution is patchy (arrowheads in D), or appears as a line corresponding to muscle ends (arrows in D,G,H,I). (J) Embryo ectopically expressing LRT-GFP (green) in muscles (red) showing its accumulation at the muscle ends (arrow) or in intracellular vesicles (arrowhead). (K) Embryo expressing LRT-GFP (green) in tendon cells. Muscles are stained with an anti-MHC antibody (red). (Inset) Arrowhead indicates membrane association of LRT in two highly magnified tendon cells. Scale bars: 20 μm in A,F,I,K.

surfaces of the tendon cells in regions where muscle ends are observed (Fig. 2D-I). In addition to the tendon-specific staining of LRT, we detected small dots along the entire body of the embryo. These dots represent non-specific staining as they also appeared in embryos homozygous for Df(2R)BSC403, which uncovers LRT, whereas the tendon-specific LRT staining was eliminated in this mutant (Fig. 1H).

In addition, we created an LRT-GFP fusion construct (in which the GFP is fused to the cytoplasmic tail of LRT), and expressed it in tendons as well as in muscle cells. In tendons, the LRT-GFP fusion protein exhibited clear surface staining (in addition to a patchy intracellular expression pattern) of the tendon cells (Fig. 2K, inset, arrowhead). When expressed in muscles, LRT-GFP accumulated at the muscle-tendon junction site in addition to small dots presumably representing vesicle staining (Fig. 2J, arrow and arrowhead correspondingly). This suggests that LRT-GFP is exported and capable of accumulating in the muscle ends. These results indicate that LRT is a tendon-specific protein that is positively regulated by Stripe and is localized at the surface of the tendon cell membrane facing muscle ends.

LRT is required for formation of the correct embryonic somatic muscle pattern

To elucidate the function of LRT, we analyzed the phenotype of embryos homozygous for a deficiency that uncovers lrt together with an additional 17 genes: Df(2R)BSC403 (for details of the other 16 genes see Fig. S1 in the supplementary material). The overall muscle pattern in these embryos appeared normal. However, we detected a specific phenotype in the ventral oblique muscles, whereby in 75% (n=15) of the mutant embryos two to four segments showed misguided extension of ventral oblique muscle 4 (VO4/muscle 31) and ventral acute muscle 3 (VA3/muscle 18) (arrowheads in Fig. 3B), whereas only 14% of wild-type embryos showed abnormal VO4/VA3 extension in a single segment (n=42) (see also Table 1). In addition, in mutant flies, thin cellular extensions were detected at the edges of most of the ventral muscles (arrows in Fig. 3B). Significantly, both phenotypes were rescued in 100% (n=11) of the mutant embryos following tendon-specific expression of full-length LRT-GFP (Fig. 3C), indicating that both muscle phenotypes stem from the lack of LRT. In addition, similar phenotypes were also detected in 73% (n=15) of embryos in which LRT was knocked down by driving the expression of RNAi against lrt together with Dicer2, using the actin-gal4 driver (Fig. 3D). The ubiquitous actin-gal4 driver was used to enable expression of the RNAi construct at early embryonic stages to reduce the mRNA levels of lrt detected at stage 12. In cells not subjected to RNAi knockdown, expressing only the actin-gal4 driver together with Dicer2, only 20% of the embryos showed abnormal VO4/VA3 extension in a single segment (n=16) (see also Table 1). Taken together, these experiments suggested that LRT is essential for the targeting of some of the ventral muscles towards their attachment sites and for the arrest of ectopic muscle extensions once the muscle reaches its corresponding tendon cell. Staining for PSβ-integrin showed that the myotendinous junction was established in the lrt mutant embryos in regions where muscles reached the tendon sites (arrows in Fig. 3E,F). However, in the ventral regions where muscles were missing, no integrin staining was detected (Fig. 3F, arrowhead), suggesting that lack of LRT does not directly affect the ability of the muscles to establish contacts with the tendon cells. Importantly, tendon cell markers, including Stripe and Shortstop were normally expressed in Df(2R)BSC403, indicating that tendon gene expression is not affected by non-related genes uncovered by the deficiency (Fig. 3G,H,I,J).

To further characterize the phenotype of the mutant embryos, we followed the morphology, migration and adhesion of a single muscle in lrt mutant embryos by recombining UAS-CD8GFP with the 5053-gal4 driver (expressed in muscle 12) on the background of Df(2R)BSC403. Whereas the majority of muscles 12 myotubes interacted normally with their corresponding tendon cells at the segmental border, we detected abnormally oriented membrane extensions of various sizes formed by this muscle (Fig. 4C,D arrows). These extensions were detected in 53% (n=23) of mutant embryos at stage 16 in at least one to two segments. Only embryos heterozygous for the 5053-gal4 driver were selected for this analysis as homozygous embryos carrying 5053-gal4 exhibit an aberrant muscle pattern. No heterozygous embryos exhibited the unique
phenotype of abnormal muscle extensions \( (n=11) \) (Fig. 4A,B). Examination of embryos at an earlier developmental stage in which muscle 12 had not arrived at its corresponding tendon cell revealed that some of these muscles exhibited extra filopodia (Fig. 4F, arrows). Significantly, the phenotype of abnormal muscle extensions was rescued in 100% of the embryos \( (n=15) \) by the expression of LRT-GFP in tendon cells, indicating that the phenotype resulted from the lack of LRT (Fig. 4G,H) (see Table 1).

Thus, the lack of LRT leads to a failure of certain muscles to locate their proper tendon cells, and in addition, causes muscles to form misguided ectopic membrane extensions even before muscle arrival at its target tendon cell.

**Overexpression of LRT changes the directional migration of certain muscles**

To further elucidate the function of LRT, we overexpressed the LRT-GFP fusion protein in several different tissues. Although ectopic expression of LRT in muscles (using the *mef2-gal4* driver) did not change muscle migration behavior, its overexpression in tendon cells led to a clear phenotype in which in 70% of the embryos \( (n=12) \) the dorsal longitudinal muscles stalled close to a single segmental border and did not extend between two segmental borders in two to four segments (Fig. 5D, arrow). In 20% of these embryos, we also detected similar defects in the ventral longitudinal muscles (Fig. 5C, arrow). In wild-type embryos, both dorsal and ventral longitudinal muscles form their initial contact with tendon cells at one side of the segment and then extend in the opposite direction, often to a tendon cell located at the more anterior segmental border. It appeared that overexpression of LRT led to stalling of the muscle in the proximity of the initial segmental border, with the muscle presumably unable to extend to the more anterior attachment site. Embryos heterozygous for *sr-gal4* did not show the ventral stalling muscle phenotype \( (n=10) \), and only 8% showed a dorsal stalling muscle phenotype \( (n=11) \) (see Table 1).

The stalled muscles often did not establish integrin-mediated adhesion with tendon cells, and the characteristic continuous line of integrin staining along the segmental border was missing or interrupted in LRT-overexpressing embryos (Fig. 5G, arrowhead). This suggests that the stalled muscles do not establish normal adhesion with their corresponding tendon cells.

In conclusion, elevating the levels of LRT in tendon cells prevented proper extension of certain muscles towards their attachment sites, inhibiting their ability to stretch between two segmental borders.

**LRT forms a protein complex with Robo receptors**

The non-autonomous effect of LRT on muscle migration behavior suggested that muscles might express a putative receptor for LRT. The leucine-rich repeat domain of LRT is similar to the second LRR domain of Slit. This domain in Slit was demonstrated to interact with Robo receptors (Hohenester, 2008; Howitt et al., 2004; Morlot et al., 2007). Importantly, the muscle phenotype of Slit and Robo mutant embryos resembles that of LRT overexpression (Kramer et al., 2001) (Fig. 7C). We therefore hypothesized that LRT might interact with Robo through its LRR domain. To test this possibility, S2 cells were first transfected with LRT-GFP alone. The GFP labeling appeared to be associated with the plasma membrane (Fig. 6A). We noticed that part of the LRT-GFP was retained inside the S2 cell, presumably representing vesicle and/or ER staining. Co-transfection with both Robo-HA and full-length LRT-GFP indicated that LRT-GFP colocalized with Robo-HA at the surfaces of the plasma membrane of the S2 cells (Fig. 6B,C,D). To verify that LRT-GFP was indeed localized at the plasma membrane, we treated live S2 cells transfected with LRT-GFP with Trypsin; the protease activity was then stopped by the addition of serum, and we analyzed the cell extract by western analysis for GFP. The levels of the intact LRT-GFP band (molecular weight =130 kDa) were significantly reduced and a new 30 kDa band appeared, corresponding in size to the truncated cytoplasmic LRT-GFP domain (Fig. 6J). This experiment verifies that LRT is indeed a transmembrane protein.

Next, a soluble form of LRT-GFP, including the full extracellular domain of LRT fused to GFP, was produced (LRT-NTD) and transfected into S2 cells. The conditioned medium of these cells, containing the secreted LRT-GFP protein, was collected and cleared from floating cells by centrifugation. It was then added to S2 cells...
transfected with Robo-HA. Strikingly, LRT-GFP accumulated specifically on the surfaces of the Robo-HA expressing cells, but not on non-transfected S2 cells (Fig. 6E-H). This experiment suggested that LRT associates specifically with Robo on the plasma membrane of S2 cells. Moreover, co-immunoprecipitation of LRT-GFP and the 5053-gal4 driver, led to an abnormal muscle pattern in which the muscles failed to extend at either end to the segmental border tendon cells. This phenotype resembled that of the LRT overexpression phenotype described above (see Fig. 5) and was detected in 47% (n=15) of the mutant embryos (Fig. 7C). Importantly, when co-expressing LRT-GFP together with the Robo RNAi constructs in the muscles, the muscle phenotype was detected in only 11% of the embryos (n=15) (Fig. 7D), suggesting that LRT potentiates Robo activity when expressed in muscles. Of note is that overexpressing LRT-GFP alone in muscles did not promote proper muscle-tendon assembly by arresting muscle filopodia formation following the arrival of the muscle at the tendon cell. This activity is presumably achieved by an interaction between LRT and Robo receptor(s) and pointing to a functional relationship between LRT and Robo.

**DISCUSSION**

When migrating cells encounter their target tissue they must arrest their migration behavior. Signals promoting arrest of cell migration are yet to be elucidated. Here we describe the identification of a novel leucine-rich repeat protein, LRT, expressed specifically by tendon cells downstream of Stripe activity, and capable of associating with muscle-expressed Robo receptors. We further demonstrated that LRT acts non-autonomously on muscles to promote proper muscle-tendon assembly by arresting muscle filopodia formation following the arrival of the muscle at the tendon cell. This activity is presumably achieved by an interaction between LRT on the tendon cell and Robo receptors on the migrating muscle cell.
LRT protein was detected at relatively late stages of embryonic development following muscle attachment to tendon cells. This suggests that it does not function primarily to guide muscles to their attachment sites, but rather is required for the arrest of muscle extension and/or migration behavior. Two possible mechanisms may account for the arrest of muscle migration behavior: a specific signal provided by the tendon cell promoting the end of filopodia formation, or alternatively the establishment of the myotendinous junction, which alters the leading edge morphology of the muscle cell so that it is no longer forming filopodia. Lack of LRT led to ectopic membrane extensions in various ventral oblique and acute muscles as well as in muscle 12, whereas in some muscles (e.g. VO4 or VA3), the continuous formation of muscle extensions in mutant embryos led to a failure to form proper muscle-tendon contacts. In other muscles (e.g. muscle 12) these extra-muscle extensions did not affect contact formation with the tendon cells. This latter phenotype supports a model by which a contact-independent signal promotes the arrest of muscle migration behavior. The expression of LRT on the surface of the tendon cell might represent such a signal.

LRT may mediate the arrest of these extraneous membrane extensions by interacting with a receptor on the muscle, which, upon binding, represses the formation of further membrane extensions. Alternatively, LRT may compete for binding to a common receptor with an attractive signal provided by the tendon cell. Association of LRT with such a receptor may dampen further signaling required for the continuous extension of the muscle leading edge. In both scenarios, overexpression of LRT in tendon cells is likely to result in a stronger association of the muscle with the proximal tendon cell, inhibiting muscle extension to the more distal tendon by arresting its filopodia formation. This phenotype was indeed observed when LRT was overexpressed in tendon cells. The muscles with extension that was arrested failed to form a proper myotendinous junction.
A possible mechanism by which LRT might act is through its interaction with the Robo receptors. Robo mediates attractive activity between the muscles and tendon cells (Kramer et al., 2001). We demonstrated here that LRT associates with the extracellular domain of Robo. In addition, we detected a reciprocal interaction between LRT and Robo, in which reducing Robo levels could rescue the muscle-stalling phenotype of LRT overexpression. This experiment suggests that LRT promotes muscle stalling through an interaction with Robo receptors. In addition, overexpression of LRT in muscles partially rescued the Robo knockdown phenotype, which is consistent with LRT potentiating Robo activity when co-expressed in the muscle. We further tested whether LRT affects the distribution of Robo receptors along the muscle membrane; however, we did not detect a major difference between Robo distribution or expression levels in Df(2R)BSC403, LRT-overexpressing, or wild-type embryos (not shown). Alternatively, LRT may inhibit ectopic membrane extensions by reducing the levels of free Robo receptors by either direct interaction with these receptors, or by association with additional muscle surface proteins required for Robo-Slit signaling.

If LRT inhibits Robo-Slit interactions, its expression at the ventral midline should inhibit muscle repulsion from this domain. Ectopic expression of LRT-GFP using the midline driver sr-gal4 slightly affected the pattern of the ventral muscles, and these muscles appeared to extend closer to the midline (see Fig. S2 in the supplementary material), suggesting that LRT counteracts Robo activity. LRT may act by modulating Robo-Slit interactions, which are essential for the correct guidance of muscles to their specific tendon-mediated insertion sites.

**Acknowledgements**

We thank F. Schnorrer, B. Dickson, S. Kramer, P. Fisher, the Bloomingpton Stock Center and the Hybridoma Center for various constructs and fly lines. We thank E. Schejter, A. Salzberg, E. Zelzer and S. Schwarzbaum for fruitful discussions and critical reading of the manuscript. This study was supported by a grant from the Israel Science Fund (ISF).

**Supplementary material**

Supplementary material for this article is available at http://dev.biologists.org/cgi/content/full/136/21/3607/DC1

**References**


