Cholinergic neurons mediate CaMKII-dependent enhancement of courtship suppression

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In Drosophila, calcium/calmodulin-dependent protein kinase II (CaMKII) activity is crucial in associative courtship conditioning for both memory formation and suppression of courtship during training with a mated female. We have previously shown that increasing levels of constitutively active CaMKII, but not calcium-dependent CaMKII, in a subset of neurons can decrease the initial level of courtship and enhance the rate of suppression of courtship in response to a mated female. In this study, we demonstrate that a subpopulation of noncholinergic, nondopaminergic, non-GABAergic neurons can cause CaMKII-dependent reductions in initial courtship, but only cholinergic neurons enhance training-dependent suppression. These data suggest that processing of pheromonal signals in two subpopulations of neurons, likely antennal lobe projection neurons, is critical for behavioral plasticity.

Modification of behavior on short time scales requires fast, likely cell signaling, changes in neuronal circuits. One molecule that has been shown in many systems to be responsible for both short-term and long-term changes in neuronal activity is calcium/calmodulin-dependent protein kinase II (CaMKII). This kinase can act a molecular switch, becoming calcium independent after autophosphorylation at T287. Production of constitutively active kinase is believed to be a first step toward establishing short- and long-term changes in neuronal properties that underlie learning (for review, see Lisman et al. 2002).

In Drosophila courtship conditioning, CaMKII is important for behavioral changes that occur during the training period and for formation of associative memory (Griffith et al. 1993; Joiner and Griffith 1997, 1999). In this behavioral paradigm, a male is exposed to a previously mated female for 1 h, and the effects of training are assessed by measuring courtship of a subsequently presented virgin female. Naive males exposed to virgin females sense female-specific stimulatory pheromones and rapidly initiate courtship, usually copulating within 15 min. In comparison, trained males show reduced courtship of virgins, and this is believed to be the result of association of stimulatory pheromones with an aversive substance given off by mated females (Tompkins et al. 1983). Memory formation can be blocked by inhibition of CaMKII in mushroom bodies, central complex, and parts of the lateral protocerebrum (Joiner and Griffith 1999). The behavior of males during training is also plastic; they initiate vigorous courtship, but over the course of training they decrease their intensity of courtship and rarely copulate. The suppression of courtship during training is driven by a neuronal circuit distinct from that mediating associative memory formation, involving neurons in the antennal lobes and lateral protocerebrum (Joiner and Griffith 1999).

The requirement for CaMKII activity was determined using overexpression of T287A and T287D CaMKII point mutants (Mehren and Griffith 2004). Expression of the constitutively active T287D CaMKII enhanced the suppression of courtship during training but had no effect on memory formation. Addition of T287A CaMKII, which is incapable of becoming calcium independent, had no behavioral effects. The effect of T287D CaMKII on courtship suppression was only seen in animals that expressed T287D in subsets of the adult brain defined by the 30YGAL4 pattern, which includes the mushroom bodies, lateral protocerebrum, antennal lobes, subesophageal ganglion, and optic lobes. These results suggested that the decision to court was not sensitive to the total amount of calcium-stimulable kinase but rather to some threshold level of constitutively active CaMKII.

In this study, we demonstrate that both initial and training-dependent changes in courtship respond to the level of calcium-independent CaMKII in a dose-dependent manner. We further show that initial courtship levels can be modulated by noncholinergic neurons, whereas the effects of T287D on training-dependent changes in courtship are mediated by cholinergic antennal lobe neurons.

Results and Discussion

To determine whether courtship suppression during training was sensitive to the absolute level of constitutively active CaMKII, we expressed T287D CaMKII under control of a tripartite GAL4/UAS/tet-off driver system (Mehren and Griffith 2004). This system allows for both spatial and temporal control of the level of transgene expression. The tetracycline transactivator (UAS-tTA) is driven by GAL4 (a yeast transcription factor that binds to the UAS element and drives transcription of UAS-linked genes) in a spatially restricted pattern. The spatial restriction of GAL4 is achieved by expressing it under control of particular defined or endogenous promoter sequences. The effector transgene (tetO-T287D) in this scheme is expressed in the GAL4 pattern, but its levels can be controlled by feeding the animal tetracycline, which inhibits transcription by tTA. This system allows complete suppression of toxic transgenes during development by growing larvae on 10 µg/mL tetracycline in the food agar. In adults, complete suppression can be achieved by feeding 100 µg/mL tetracycline in sucrose. Removal or reduction of the drug leads to induction of the transgene.

Previous work had shown that expressing T287D with tetracycline-controlled 30YGAL4 and allowing maximal adult in-
uction by placing newly eclosed males on food containing no tetracycline was able to cause both a lowered level of initial courtship and an enhancement of trainer-dependent suppression (Mehren and Griffith 2004). If the mechanism of calcium-independent CaMKII’s actions was dependent on the level of calcium-independent CaMKII, we would expect that manipulating the amount of induction by placing newly eclosed males on a range of drug concentrations from 100 to 0 µg/mL should produce a dose-dependent range of levels of courtship suppression. Figure 1 shows that after rearing on a maximally suppressing dose of tetracycline, suppression scales with the adult level of tetracycline. Initial courtship (Fig. 1A) is decreased with increasing expression of T287D and the animals fall into two groups that are statistically different: Untreated wild-type males, fully suppressed (100 µg/mL tetracycline) T287D males, and T287D males maintained on 10 µg/mL tetracycline all show high initial courtship, whereas animals on 1 or 0 µg/mL tetracycline have a similar, low level of courtship. This suggests that there is a bimodal relationship between the level of constitutive CaMKII and initiation of courtship.

The drop in courtship caused by continued exposure to the mated female is also sensitive to the level of CaMKII. Males with additional T287D CaMKII begin to suppress courtship immediately without the characteristic lag seen in wild-type animals (Fig. 1B) (Mehren and Griffith 2004). But in contrast to the effects of T287D dosage on initial courtship, the effects on the trainer-specific component are more graded over the range of doses administered. Animals expressing the maximum level of T287D (fed 0 µg/mL tetracycline as adults) suppress courtship toward the mated female (CI/CI₀ = 0.04 ± 0.04, data presented as mean ± SEM) 12-fold better than animals fed 10 µg/mL tetracycline (CI/CI₀ = 0.47 ± 0.14), and 18-fold better than fully suppressed animals (fed 100 µg/mL tetracycline [CI/CI₀ = 0.70 ± 0.09]). The difference in the effects of particular doses of T287D on initial and trainer-dependent suppression suggests that these effects may be mediated by distinct sets of cells that innervate the antennal lobe.

To refine the mapping of these two types of courtship suppression, we utilized GAL4 lines that are restricted to neurons expressing particular neurotransmitters. The antennal lobe is known to contain synapses made by cholinergic olfactory receptor neurons (Barber et al. 1989), cholinergic projection neurons (Yasuyama et al. 2003), GABAergic interneurons (Ng et al. 2002; Wilson and Laurent 2005) and perhaps other neurons that release modulatory peptides or amines (Python and Stocker 2002). By restricting T287D expression to adult neurons of particular neurochemical classes, we can subdivide the antennal lobe circuit. Figure 2A shows the expression patterns of Cha-GAL4 (restricted to cholinergic neurons), TH-GAL4 (restricted to dopaminergic neurons), GAD-GAL4 (restricted to GABAergic neurons) as assessed by mCD8–GFP expression. All of these neurochemical GAL4s show antennal lobe expression.

We can also neurochemically subdivide the 30Y-GAL4 circuitry using GAL80 (Lee and Luo 1999). GAL80 is a yeast protein that antagonizes GAL4 action. Expression of GAL80 under control of the Cha promoter will block GAL4 action exclusively in the cholinergic subpopulation of a particular GAL4 pattern (Kimatelyo 2002). Figure 2B shows the effect of adding Cha-GAL80 to 30Y-GAL4. All antennal lobe expression of UAS-mCD8–GFP is lost, but substantial expression remains in the mushroom body, pars intercerebralis, and extrinsic mushroom body neurons, indicating that these neurons are not cholinergic.

To assess the contribution of these neurons to the T287D-dependent decrease in initial courtship, we examined the ratio of courtship during the first 10 min of exposure to a mated female (CI/CI₀) of males with adult expression of constitutive CaMKII in neurochemical subsets of cells to the mean of their GAL4 driver control (Fig. 3A). CI₀ for effector transgene controls (tetO-T287D/+; UAS-tTA+/+, 0.51 ± 0.05; UAS-tTA+/+; Cha-GAL80/+, 0.68 ± 0.05) did not differ significantly from those of wild type (0.60 ± 0.05) (data presented as mean ± SEM and analyzed by ANOVA F[2,64] = 2.78, P = 0.07). A ratio of less than one indicates a T287D-specific decrease in initial courtship and controls for nonspecific effects of individual GAL4 drivers. Consistent with our observations in Figure 1A, initial levels of courtship in animals expressing T287D under control of 30Y-GAL4 are significantly lower than the GAL4 control. Interestingly, TH-GAL4, Cha-GAL4, and GAD-GAL4 drivers cannot cause a T287D-specific decrease in initial courtship. In agreement with this, removal of the cholinergic component of the 30Y pattern by coexpression of Cha-GAL80 does not alter the effects of T287D, indicating that the cells marked by 30Y that are controlling initial courtship are noncholinergic.

Training-dependent courtship suppression can also be dissected neurochemically by looking at the training index (CI/CI₀) normalized to the training index of the GAL4 control. Again, a ratio of less than one indicates a T287D-specific effect on trainer-dependent suppression. Animals expressing T287D under
and innervate the antennal glomeruli. Since T287D expression in primary sensory neurons does not produce an enhancement of training (Mehren and Griffith 2004), the most likely antennal cholinergic neurons to be responsible for this plasticity effect are the projection neurons. We have also previously shown that 30Y-GAL4 does not express in antennae (Mehren and Griffith 2004) where primary olfactory neurons reside, supporting the idea that the CaMKII-sensitive cholinergic neurons of the 30Y-GAL4 pattern are projection neurons. This makes it likely that projection neurons are active participants in CaMKII-dependent short-term plasticity of courtship behavior.

**Materials and Methods**

**Fly lines**

Fly stocks and crosses were maintained on cornmeal, yeast, dextrose, and agar medium and kept at 25°C in a 12-h light/12-h dark cycle. Tetracycline feeding was performed by growing larvae on medium containing tetracycline at 10 µg/mL food or by feeding tetracycline to adults at a concentration of 0, 1, 10, or 100 µg/mL in a 4% sucrose solution, as described in Mehren and Griffith (2004). Canton S flies were used as wild type, and all transgenic lines were in a Canton S, white background. UAS-tTA was a gift from Bruno Bello (University of Basel, Basel, Switzerland). The tetO-CaMKII-T287D line was described in Mehren and Griffith (2004). The lines 30Y-GAL4 (Yang et al. 1995), Cha-GAL4 (Salvatore and Kitamoto 2001), Cha-GAL80 (Kitamoto 2002) and TH4B-GAL4 (Friggi-Grelin et al. 2003) have been previously described.

**Immunohistochemistry**

Brains were dissected in PBS from 5- or 6-d-old adult flies, fixed in 4% paraformaldehyde for 20–30 min, washed in PBS, and mounted in glycerol and Vectashield (Vector Laboratories Inc.). Images were acquired using the 40× objective on a Leica TCS SP2 confocal scanning microscope.

**Behavior**

All behavioral observations were performed at 25°C and 70% relative humidity in a Harris Environmental Room, in dim red light, as described in Mehren and Griffith (2004). Courtship conditioning was carried out by placing individual males with fertilized females (mated the day before) in Plexiglas mating chambers (8 mm diameter, 3 mm high) for 1 h. Courtship indices (CIs) were measured for 3 time intervals during training (0–10 min [Cl$_{10}$], 20–30 min, and 50–60 min [Cl$_{60}$]). A training index was taken as the fraction of the Cl$_{10}$ over the Cl$_{60}$. Flies who had Cl$_{60}$<0.1 were considered “nonbehaving” and therefore were excluded from analysis. Each CI was subjected to arcsine square root transformation to approximate normal distributions. Transformed CIs or training indices were subjected to a one-way ANOVA with genotype as the main effect, using JMP (version 5.0.1.2; SAS Institute), and planned comparisons of means were performed with Student’s t-test (adjusted for experiment-wise error).
Figure 3. Neurochemical restriction separates initial and trainer-dependent effects of T287D CaMKII. In all experiments T287D CaMKII expression was limited to adult neurons by use of the GAL4/UAS/tet-off system with 10 µg/mL tetracycline during development and no tetracycline feeding in the adult stage. (A) Initial courtship levels are modulated by expression of T287D CaMKII in noncholinergic cells. C_l0 data for T287D CaMKII-expressing males was normalized to the mean of the C_l0 for the appropriate GAL4 line. A normalized ratio of <1.0 means that T287D CaMKII reduced initial courtship. Flies expressing 30Y- or 30Y/Cha-GAL80-driven T287D CaMKII show significantly less initial courtship of mated females than do GAL4 controls (ANOVA F[8,147] = 3.37, P = 0.0014, with post hoc comparisons significantly different, α = 0.01). (B) Cholinergic cells mediate T287D-dependent enhancement of courtship training. Training index (C_lf/C_l0) data for T287D CaMKII-expressing males was normalized to the mean of the training index for the appropriate GAL4 line. A normalized ratio of <1.0 means that T287D CaMKII enhanced training-dependent courtship suppression. When 30Y-GAL4 is used to drive T287D CaMKII expression in adults only, training performance is enhanced, however when Cha-GAL80 is used to inhibit expression of T287D in cholinergic cells of the 30Y-GAL4 pattern, training enhancement is rescued. When T287D is expressed solely in cholinergic cells using Cha-GAL4, training enhancement is similar to enhancement with 30Y-driven T287D. CaMKII expression in dopaminergic or GABAergic cells in the adult brain does not significantly affect courtship training with a mated female. ANOVA F[8,146] = 5.03, P < 0.0001, with post hoc comparisons between 30Y-GAL4/T287D or Cha-GAL4/T287D, and GAL4 controls significantly different, α = 0.01. n = 9–22.

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