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Ca$_{\text{V}3}$-type $\alpha$1T calcium channels mediate transient calcium currents that regulate repetitive firing in Drosophila antennal lobe PNs

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Iniguez J, Schutte SS, O'Dowd DK. Ca$_{\text{V}3}$-type $\alpha$1T calcium channels mediate transient calcium currents that regulate repetitive firing in Drosophila antennal lobe PNs. J Neurophysiol 110: 1490–1496, 2013.

A WIDE VARIETY OF INSECT BEHAVIORS are driven or modulated by olfactory input and the ensemble of neurons involved in processing olfactory information is well defined (Wilson and Mainen 2006; Voßhall and Stocker 2007). Olfactory perception begins when odorant molecules bind to receptors in olfactory receptor neurons (ORNs) located in the antennae and the maxillary palps (Hildebrand and Shepherd 1997). ORNs project to the antennal lobes, the insect equivalent of the vertebrate olfactory bulb, where they synapse onto the dendrites of projection neurons (PNs), the principal output cells that extend axons to higher order processing centers in the mushroom bodies and lateral horn (Stocker 1994; Ito et al. 1998). In Drosophila melanogaster, where genetic manipulations and behavioral assessment are routine, it is now feasible to record from identified neurons within this circuit in the adult brain (Wilson et al. 2004; Gu and O'Dowd 2006, 2007). This has made it possible to explore the mechanisms that regulate activity in a circuit important in generating specific components of an adult behavior at the single cell level. Whole cell recordings from single PNs have demonstrated that olfactory processing begins in the antennal lobe where both intra- and interglomerular interactions influence activity of these cells (Olsen et al. 2007; Olsen and Wilson 2008; Root et al. 2008).

To understand the molecular mechanisms underlying regulation of neuronal activity in individual PNs in the olfactory circuit requires identification of the ion channel subtypes that govern excitability and synaptic transmission in these cells. One important class of channels present in all neurons are voltage-gated calcium channels that mediate depolarization-induced calcium influx that influences a number of cellular processes including excitability and release of neurotransmitters at chemical synapses. The $\alpha$1-subunit of these multimeric proteins forms the ion-conducting pore that defines many of the functional properties characteristic of the distinct calcium channel subtypes (Catterall 2011). There are three families of genes encoding $\alpha$1 subunits and in the Drosophila genome there is one $\alpha$1 subunit gene in each family: $\alpha$ID (Ca$_{\text{V}1}$), cac (Ca$_{\text{V}2}$), and $\alpha$1T (Ca$_{\text{V}3}$) (Zheng et al. 1995; Smith et al. 1996; Littleton and Ganetzky 2000; King 2007). In a recent study we found that voltage-gated calcium currents recorded from the cell bodies of PNs in the adult brain could be separated into two kinetically distinct components: a rapidly decaying transient current and a slowly decaying sustained current (Gu et al. 2009). Using a combination of pharmacological and molecular genetic strategies, we demonstrated that the Ca$_{\text{V}2}$-type cac gene encodes calcium channels that mediate PLTXII-sensitive sustained calcium currents. Our studies show that the CAC channels regulate action potential-independent release of neurotransmitter at excitatory cholinergic synapses in the adult brain, a novel role not predicted from previous studies at peripheral synapses (Rieckhoff et al. 2003).

While a recent study suggests that Ca$_{\text{V}2}$-type CAC channels (aka Dmca1A) also contribute to the transient calcium currents in adult motor neurons (Ryglewski et al. 2012), our results indicated that neither PLTXII nor mutations in the cac gene reduced the transient currents in PNs. This suggests that a distinct calcium channel subtype gives rise to the transient current in adult PNs. In vertebrates, previous studies have demonstrated that Ca$_{\text{V}3}$ genes encode channels underlying transient calcium currents (Nowycky et al. 1985; Catterall 2011).

Here we report that amiloride, a vertebrate Ca$_{\text{V}3}$-type channel blocker, reduces the transient calcium current without significantly altering the sustained Ca$_{\text{V}2}$-type CAC channel-mediated current in adult PNs. In addition, RNAi mediated knockdown of the $\alpha$1T gene, the Drosophila Ca$_{\text{V}3}$-type homolog, in PNs reduced the transient component significantly.
but the sustained component was not affected. Alterations in evoked and spontaneous firing were observed in the aIT knockdowns. These data demonstrate that aIT-encoded Ca$_{3,3}$-type channels mediate transient calcium currents that are important in shaping the PN firing properties and, therefore, play an important role in regulating olfactory signal processing.

**METHODS AND MATERIALS**

*Fly strains.* All recordings of wild-type PNs were from the GH146-Gal4 line. To knockdown expression of channels encoded by the *Drosophila* Ca$_{3,3}$-type calcium channel gene (aIT) specifically in PNs, the GH146-Gal4 line was crossed to two independent UAS-RNAi lines from Vienna Drosophila RNAi Center targeting different sequences in the aIT gene. The F1 fly lines were designated RNAi-1 Ta (no. 48008) and RNAi-1 Tb (no. 31961), respectively. All fly lines were kept at 23°C in standard plastic vials with cotton plugs on a yeast, agar, and cornmeal diet with a 12 h light/dark regimen.

Electrophysiological recordings from PNs in isolated adult brain. Brains were obtained from adult female flies 1–2 days after eclosion. The entire brain, including optic lobes, was removed from the head, prepared for recordings as previously described, and mounted in the recording chamber with the anterior face of the brain up (Gu and O’Dowd 2006, 2007). Recordings were made from PNs in the dorsal neuron cluster using 8- to 9-mΩ resistance pipettes.

Isolated calcium currents were recorded in voltage-clamp mode using a pipette solution containing: 102 mM d-gluconic acid, 102 mM CsOH, 0.085 mM CaCl$_2$, 1.7 mM MgCl$_2$, 17 mM NaCl, 0.94 mM EGTA, 8.5 mM HEPES, and 4 mM Na$_2$ATP. The osmolarity was adjusted to 235 mosM and pH to 7.2. The external solution contained: 101 mM NaCl, 1.8 mM CaCl$_2$, 0.8 mM MgCl$_2$, 5.4 mM KCl, 5 mM glucose, 1.25 mM NaH$_2$PO$_4$, and 20.7 mM NaHCO$_3$. Drugs were added to block synaptic transmission, sodium current, and potassium currents: 20 μM t-tubocurarine (Sigma), 10 μM picrotoxin (Sigma), 1 mM tetrodotoxin (Alomone), 2.5 mM TEA (Sigma), and 1 mM 4-aminopyridine (Sigma). The osmolarity was adjusted to 250 mosM, and the solution was saturated with 95% oxygen-5% CO$_2$. For experiments evaluating the effect of amiloride (Sigma), the drug was made up fresh each day by dissolving in H$_2$O. This was added to the extracellular solution for a final concentration of 1 mM.

Firing properties were recorded in current-clamp mode with internal solution as described above except that cesium gluconate was replaced by potassium gluconate. Intrinsic firing properties, both evoked and spontaneous, were evaluated in external solution that contained picrotoxin and t-tubocurarine to block synaptic transmission. All recordings were performed during continuous perfusion with oxygenated saline at room temperature.

*Data acquisition and analysis.* A Dell computer (Dimension 8200) was used in conjunction with an EPC-7 patch clamp amplifier (List Medical Electronics, Darmstadt, Germany), Digitida 1322A analog to digital converter (Axon Instruments), and Clampex software (pClamp 9.0; Molecular Devices) filtered through a 2-kHz low-pass Bessel filter (Frequency Devices) for generation of voltage/current commands and for data acquisition. Current amplitudes were normalized to whole cell capacitance, and values are reported as current density. Comparisons between two genotypes or treatments were made using an ANOVA with Bonferroni’s post hoc test for pair-wise comparisons. Data shown was corrected for the 5-mV liquid junction potential associated with the recording solutions. Statistical significance was assumed when *P* < 0.05, **P** < 0.01, and ***P*** < 0.001. Data in graphs are presented as means ± SE.

**RESULTS**

Amiloride inhibits transient calcium currents in adult PNs. Antennal lobe PNs in brains of wild-type adult flies express voltage-gated calcium currents with two kinetically distinct components. In a previous study we demonstrated that Ca$_{2,1}$-type CAC channels are major contributors to the PLTXII-sensitive sustained current based on selective reduction of this component in *cac* hypomorphic mutants (Gu et al. 2009). As the first step in identifying the channels underlying the transient calcium component we used a pharmacological approach focusing on the drug amiloride that blocks T-type currents in vertebrate neurons. Isolated calcium currents were recorded in physiological concentrations of extracellular calcium (1.8 mM) with drugs to block both sodium and potassium currents.

Depolarizing voltage-steps from a holding potential of −80 mV evoked calcium currents with both a transient and sustained component in a wild-type PN in the adult brain preparation (Fig. 1A). The time course of activation of the transient current is relatively slow with the current peak occurring measured at half amplitude. The number of spikes/burst was determined for each burst and a mean value was calculated for each cell. Analysis was conducted in PNs in which the resting membrane potential was more hyperpolarized than −40 mV and that exhibited at least one burst of action potentials during the 2.5-min recording period.

Statistics. Comparisons between two genotypes or treatments were made with Student’s t-test. Comparisons between more than two genotypes or treatments were made using an ANOVA with Bonferroni’s post hoc test for pair-wise comparisons. Data shown was corrected for the 5-mV liquid junction potential associated with the recording solutions. Statistical significance was assumed when *P* < 0.05, **P** < 0.01, and ***P*** < 0.001. Data in graphs are presented as means ± SE.

**Fig. 1.** Transient calcium currents in projections neurons (PNs) are amiloride sensitive. A: isolated calcium currents in a PN in brain of wild-type adult with both transient and sustained components. Currents elicited by a series of depolarizing voltage steps (−50, −40, −20, and −10 mV) from a holding potential of −80 mV; B: perfusion of recording chamber with 1 mM amiloride reduced the transient component with little effect on the sustained currents; C: amiloride-sensitive current isolated by digital subtraction of current records (A and B). D: current density vs. voltage curve generated from digitally generated amiloride-sensitive transient currents (*I-V; n = 5*). Bars indicate SE. Peak current at each voltage was normalized to cell capacitance and expressed as density (pA/pF). Isolated calcium currents were recorded in external saline containing physiological concentration of calcium (1.8 mM), tetrodotoxin (TTX), picrotoxin (PTX), curarine, TEA, and with internal cesium.

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Amiloride caused a significant reduction in the magnitude of the transient current in the population of PNs examined (Fig. 2C; \( P < 0.001 \), unpaired \( t \)-test). The magnitude of the sustained current was not significantly reduced by amiloride (Fig. 2D). Together these findings indicate that amiloride-sensitive calcium channels mediate transient calcium currents in PNs in adult brain.

**RNAi-1T knockdown in PNs reduces transient calcium currents.** Based on kinetics and pharmacological profile, the *Drosophila* \( \alpha \)IT gene, which is homologous to the vertebrate \( \alpha \)3-type calcium channels, was a good candidate for encoding the channels underlying the transient current in PNs. To test this hypothesis, flies carrying UAS-RNAi targeted to the \( \alpha \)IT gene were crossed to the GH146-Gal4 PN driver line with the goal of specifically knocking down \( \alpha \)IT channel expression in PNs. For these studies we used two different UAS-RNAi lines, targeting different sequences in the \( \alpha \)IT gene and representing independent insertion events on the second and third chromosome. These were designated RNAi-1T-a and RNAi-1T-b.

To determine if the RNAi knockdown lines exhibited altered calcium currents in PNs, the prepulse inactivation protocol was used to evaluate the magnitude of the transient and sustained current components. Representative records of calcium currents from both RNAi-1T knockdown lines illustrate a relatively small transient component while the sustained component remains prominent (Fig. 3, A and B). The magnitude of the transient current in RNAi-1T-a knockdown PNs was significantly reduced in RNAi-1T-a (\( n = 8 \)) and RNAi-1T-b (\( n = 9 \)) compared with wild type (\( n = 14 \); \( \ast \ast \ast P < 0.001 \), ANOVA, Bonferroni’s post hoc test). D: magnitude of sustained calcium current evaluated from same PNs were not significantly different than wild type. Whole cell calcium currents were isolated with TTX, PTX, \( \beta \)-tubocurarine, TEA, and internal cesium. Each bar indicates the means and SE.

**Fig. 3. RNAi-1T knockdown specifically reduces transient calcium currents in PNs.** A and B: prepulse inactivation protocol reveals PN calcium currents with small amplitude transient component in RNAi-1T-a and RNAi-1T-b knockdowns. The sustained component is still prominent. C: magnitude of the transient calcium current determine from prepulse protocol is significantly reduced in RNAi-1T-a (\( n = 8 \)) and RNAi-1T-b (\( n = 9 \)) compared with wild type (\( n = 14 \); \( \ast \ast \ast P < 0.001 \), ANOVA, Bonferroni’s post hoc test). D: sustained calcium current magnitude evaluated from same PNs were not significantly different than wild type. Whole cell calcium currents were isolated with TTX, PTX, \( \beta \)-tubocurarine, TEA, and internal cesium. Each bar indicates the means and SE.
RNAi-1T knockdown alters spontaneous burst firing in PNs. 1T channels in other systems are important in regulation of spontaneous, pacemaker activity in excitable cells (Catterall 2011). To determine if 1T channels are also involved in regulating spontaneous activity in PNs, 1T-tubocurarine and picrotoxin were added to the bath to block cholinergic and GABAergic synaptic transmission. In addition whole cell current clamp recordings were made in the absence of applied holding current. There was no significant difference in the mean resting potential between the two genotypes (Table 1).

In the presence of synaptic current blockers, PNs in both wild-type and RNAi-1T-a knockdown brains exhibit spontaneous membrane potential depolarizations (Fig. 5A). These vary in frequency, amplitude, and duration during continuous recording in a single cell so evaluation of spontaneous activity was based on assessment during the first 2.5 min of each recording. Bursts were defined as suprathreshold depolarization copped by one or more spikelets (Fig. 5A). Burst properties, including mean frequency, amplitude, duration, and number of spikelets/burst were evaluated for each cell. Subthreshold depolarizations were not analyzed (Fig. 5A).

While the mean spikelet frequency/burst was larger in RNAi-1T-a than wild type, the difference was not significant (Fig. 5B). However, there was a fourfold increase in the number of spikelets/burst in the RNAi-1T-a knockdown compared with wild type (Fig. 5C, $P < 0.01$, $t$-test). Therefore, we also examined the burst duration and found this was significantly longer in RNAi-1T-a than wild type (Fig. 5, A and C, $P < 0.05$, $t$-test). The increase in number of spikelets/burst is consistent with the increase in evoked firing frequency and suggests the 1T channel is important in regulating action potential frequency in spontaneously firing PNs. The increase in burst duration suggests 1T channels also contribute to the waveform of the low-frequency depolarizations that give rise to burst firing.

### Table 1. Analysis of spontaneous activity indicates no significant difference in the resting potential, burst frequency, or burst amplitude in wild-type and RNAi-1T-a knockdown PNs

<table>
<thead>
<tr>
<th></th>
<th>Resting Potential, mV</th>
<th>Burst Frequency, Hz</th>
<th>Burst Amplitude</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild type</td>
<td>$-59.5 \pm 1.47$</td>
<td>$0.22 \pm 0.1 \pm$</td>
<td>$12.2 \pm 1.7$</td>
</tr>
<tr>
<td>RNAi-1T-a</td>
<td>$-56.75 \pm 1.63$</td>
<td>$0.21 \pm 0.05 \pm$</td>
<td>$10.1 \pm 0.9$</td>
</tr>
<tr>
<td>$t$-test</td>
<td>$P = 0.22$</td>
<td>$P = 0.94$</td>
<td>$P = 0.31$</td>
</tr>
</tbody>
</table>

Values are means ± SE. PNs, projection neurons.
to spontaneous burst firing characteristic of PNs. In contrast, while the frequency and amplitude of the bursts were quite variable between PNs, there was no significant difference between the two genotypes (Table 1).

**DISCUSSION**

Projection neurons in the adult *Drosophila* antennal lobe process olfactory input from olfactory receptor neurons and relay this to neurons in the mushroom body and lateral horn. This well-defined neural pathway and access to these neurons in the whole brain preparation provide an excellent model system to explore how voltage-gated calcium channels regulate signaling in an identified neuronal subtype within the adult olfactory circuit. The results of this study demonstrate that the transient calcium current in the adult *Drosophila* antennal lobe PNs is mediated by amiloride-sensitive αIT-encoded channels. Furthermore, the αIT channels were found to modulate cell excitability, a novel role in the *Drosophila* central nervous system.

A recent analysis of adult *Drosophila* motor neurons reported three distinct calcium current phenotypes in an αIT null mutant: complete loss of all calcium currents (56%), reduction of both sustained and transient currents (28%), and no effect on calcium currents (16%) (Ryglewski et al. 2012). In contrast, when we used a PN-specific Gal4 driver in conjunction with two independently generated UAS-RNAi lines targeted to the αIT gene, this resulted in specific reductions in the transient calcium current. No significant change in the sustained calcium current was observed. Since the two RNAi lines were targeted to distinct regions of the αIT gene, this is compelling evidence that the αIT gene encodes channels underlying the transient current in PNs. The varied effects of an αIT null mutant of calcium currents previously reported in motor neurons may be associated with activation of homeostatic regulatory mechanisms caused by elimination of the channel in all cell types.

Similar to currents mediated by T-type channels in vertebrates, the transient calcium currents in adult PNs are inhibited by amiloride, a vertebrate Ca_{3,2} T-type calcium channel blocker (Tang et al. 1988; Hirano et al. 1989). Amiloride has also been reported to block a portion of the calcium current that could be inactivated by depolarization in *Drosophila* larva body wall fibers and embryo motor neurons (Gielow et al. 1995; Baines and Bate 1998). While the underlying gene encoding the current in muscle and motor neurons was not identified, our data suggest the αIT gene may also encode the channels in these cells. In a recent study it appears that αIT channels also underlie an amiloride-sensitive transient current that represents a relatively small component of the total calcium current in adult motor neurons (Ryglewski et al. 2012). The majority of transient current in the motor neurons, however, appears to be mediated by Ca_{2,2}-type CAC channels (Ryglewski et al. 2012). This suggests that, similar to finding in the mammalian central nervous system, stage and/or cell-specific splicing events give rise to calcium channels with distinct functional properties (Bell et al. 2004; Lipscombe et al. 2012).

In contrast to T-type currents in vertebrates that typically activate at low voltages, the PN transient calcium currents are first activated at a membrane potential of between −50 and −40 mV. This activation profile is similar to that reported for transient calcium currents in neurons cultured from brains of late stage wild-type pupae (Gu et al. 2009), *Drosophila* embryonic motor neurons (Byerly and Leung 1988), *Drosophila* muscle fibers (Gielow et al. 1995), and *Drosophila* larval motor neurons (Worrell and Levine 2008). In contrast, studies in cytokinesis-arrested neuroblasts (Peng and Wu 2007), embryonic motor neurons (Baines and Bate 1998), and adult motor neurons (Ryglewski et al. 2012) reported both low- and high-voltage-activated calcium currents. The discrepancies in the studies could arise from the differences in cell types and developmental stages. It is also possible that contributing to the differences noted is the use of barium vs. calcium as the charge carrier for studying calcium influx. Replacing calcium with barium substantially increases the current amplitude (Byerly and Leung 1988). Barium is also known to suppress Ca^{2+}-dependent inactivation of calcium channels. In larval motor neurons when barium was used to replace calcium as the charge carrier, this caused a substantial increase in current amplitude and altered the kinetics of calcium current decay (Worrell and Levine 2008). All of the recordings in PNs in the present study were conducted in physiological concentrations of Ca^{2+}, and therefore, calcium-dependent inactivation of the current could contribute to the transient nature of the current.

Fig. 5. αIT knockdown affects spontaneous firing in PNs. A: spontaneous activity in a wild-type and a RNAi-1T-a knockdown PN. Low-frequency, slow-wave depolarizations, classified as bursts, are marked by arrows. Sub-threshold depolarizations (sub) were not analyzed. Dashed line indicates the resting membrane potential. One burst from each PN is expanded illustrating slow-wave depolarizations, classified as bursts, are marked by arrows. Sub-threshold depolarizations (sub) were not analyzed. Dashed line indicates the resting membrane potential. One burst from each PN is expanded illustrating slow-wave depolarizations capped by rapid spikelets. The amplitude, duration, and number of spikelets were evaluated as indicated for each burst in the first 2.5 min of recording. B: mean spikelet frequency is not significantly different in wild-type and the RNAi-1T-a knockdown. C: however, mean number of spikelets/burst is significantly higher in RNAi-1T-a knockdown vs. wild-type PNs (**P < 0.01, t-test). D: Mean burst duration is significantly longer in RNAi-1T-a knockdown PNs compared with wild-type (*P < 0.05, t-test).

Whole cell current clamp recordings done in PTX and curare to block synaptic activity. Bars indicate means ± SE; n = 11 for wild-type and n = 12 for RNAi-1T-a knockdowns.
The increase in evoked firing frequency in PNs we report in the RNAi-1T knockdowns demonstrates that the α1T channels are important in regulating excitability in adult *Drosophila* neurons. The increased excitability caused by reducing expression of Ca₃,3-type channels in the adult PNs is similar to the upregulation of firing frequency documented in larval motor neurons following genetic reduction of Ca₃,1 or Ca₃,2 type calcium channels (Worrell and Levine 2008). In the wild-type motor neurons, the increase in firing frequency was mimicked by acute removal of calcium from the recording solution suggesting the change in excitability reflects reduced activation of Ca²⁺-activated K⁺ channels. However, Peng and Wu (2007) were not able to rule out the possibility that there was compensatory downregulation of other K⁺ channel genes in the Ca₃,1 and Ca₃,2 mutants that could have the same effect of increasing motor neuron-evoked firing frequency.

To avoid the possible contribution of developmental regulation of other channel types, it would be helpful to explore the role of α1T channels in regulating excitability by acutely blocking these channels in wild-type PNs. Unfortunately *Drosophila* express nonvoltage-gated sodium channels that are also sensitive to amiloride (Adams et al. 1998; Liu et al. 2003; Zelle et al. 2013), and our preliminary studies indicate these can affect membrane potential depolarizations directly (data not shown). Therefore, additional experiments with double mutant combinations will be necessary to determine if reduced activation of Ca²⁺-activated K⁺ channels and/or reduced expression of other K⁺ channels contribute to the increase in firing frequency seen in adult PNs in α1T knockdowns.

Low-voltage-activated T-type Ca²⁺ channels in mammals have been shown to be important in pacemaking activity in sino-atrial node of the heart (Mangoni et al. 2006), and they are crucial for generation of rhythmic bursts of action potentials in thalamic relay neurons of the thalamus (Perez-Reyes 2003). In PNs there was no significant change in the burst firing frequency, but the burst duration in α1T knockdowns was significantly increased. This suggests that while T-type calcium channels do have a role in regulating spontaneous burst firing in *Drosophila* PNs, their relatively high activation voltage may limit their contribution to pacemaking type activities.

Is the third calcium channel subtype encoded by the DMCAID gene also expressed in adult PNs? Our preliminary studies revealed that calcium currents examined in PNs from homozygous α1D mutant AR66 and α1D-knockdowns (VDRC no 51491) had both transient and sustained currents that were not significantly different from wild-type (data not shown). However, alteration in the firing properties of DMCAID knockdown and mutant indicates that these channels are expressed in PNs. This suggests that the currents mediated by these channels are located in the axonal compartment, electrically distant from the soma. If calcium influx is initiated at some distance from the electrode, this current change will be substantially attenuated when it reaches the soma. Unfortunately, the soma of PNs is the only location in these cells that is accessible to the patch-clamp electrode (Gouwens and Wilson 2009).

In conclusion, our results demonstrate for the first time that α1T-encoded voltage-gated calcium channels are expressed in adult PNs where they are important in regulating excitability. Further studies on how these channels regulate olfactory-related behavior will be an important contribution to our understanding of how activity within specific neurons in a well-defined circuit guides behavior.

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**DISCLOSURES**

No conflicts of interest, financial or otherwise, are declared by the author(s).

**AUTHOR CONTRIBUTIONS**


**REFERENCES**


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**DISCLOSURES**

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