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Abstract: Lymphatic vessels (LVs) are important in the regulation of tissue fluid homeostasis and the pathogenesis of tumor progression. We investigated the innervation of LVs and the response to agonists and antagonists of the autonomic nervous system *in vivo*. While skin-draining collecting LVs express muscarinic, α - and β -adrenergic receptors on lymphatic endothelial cells and smooth muscle cells, intestinal lacteals express only β -adrenergic receptors and muscarinic receptors on their smooth muscle cells. Quantitative *in vivo* near-infrared imaging of the exposed flank-collecting LV revealed that muscarinic and β -adrenergic agonists increased LV contractility, whereas activation of α -adrenergic receptors inhibited contractility and initiated nitric oxide (NO)-dependent vasodilation. Tumor-draining LVs were expanded and showed a higher innervation density and contractility that was reduced by treatment with atropine, phentolamine, and, most potently, isoproterenol. These findings likely have clinical implications given the impact of lymphatic fluid drainage on intratumoral fluid pressure and thus drug delivery.

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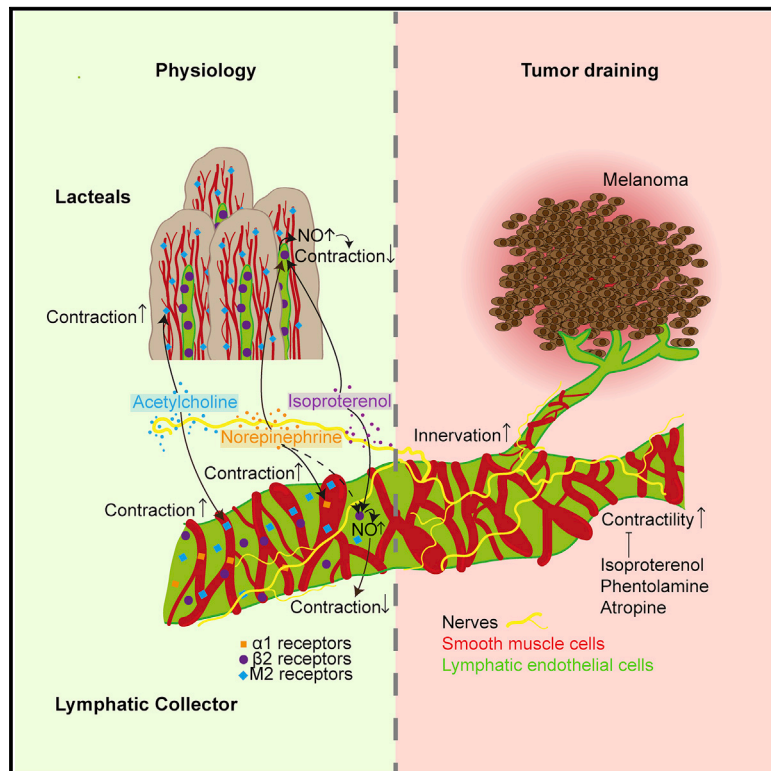
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A Distinct Role of the Autonomic Nervous System in Modulating the Function of Lymphatic Vessels under Physiological and Tumor-Draining Conditions

Graphical Abstract



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In Brief

Bachmann et al. use *in vivo* imaging to demonstrate that muscarinic and α_1 -adrenergic agonists increase lymphatic contractility, whereas activation of β_2 -adrenergic receptors inhibits contractility and mediates NO-dependent vasodilation. Tumor-draining lymphatics have a denser innervation and increased contractility, which can be diminished by β_2 -adrenergic agonists.

Highlights

- Murine lymphatic vessels are innervated in an organ-specific manner
- α_1 -adrenergic and muscarinic agents enhance lymphatic contractility *in vivo*
- β_2 -adrenergic agents reduce lymphatic contractility
- Tumor-draining lymphatic vessels have increased innervation and contractility



A Distinct Role of the Autonomic Nervous System in Modulating the Function of Lymphatic Vessels under Physiological and Tumor-Draining Conditions

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SUMMARY

Lymphatic vessels (LVs) are important in the regulation of tissue fluid homeostasis and the pathogenesis of tumor progression. We investigated the innervation of LVs and the response to agonists and antagonists of the autonomic nervous system *in vivo*. While skin-draining collecting LVs express muscarinic, α_1 - and β_2 -adrenergic receptors on lymphatic endothelial cells and smooth muscle cells, intestinal lacteals express only β -adrenergic receptors and muscarinic receptors on their smooth muscle cells. Quantitative *in vivo* near-infrared imaging of the exposed flank-collecting LV revealed that muscarinic and α_1 -adrenergic agonists increased LV contractility, whereas activation of β_2 -adrenergic receptors inhibited contractility and initiated nitric oxide (NO)-dependent vasodilation. Tumor-draining LVs were expanded and showed a higher innervation density and contractility that was reduced by treatment with atropine, phentolamine, and, most potently, isoproterenol. These findings likely have clinical implications given the impact of lymphatic fluid drainage on intratumoral fluid pressure and thus drug delivery.

INTRODUCTION

The lymphatic vascular system is responsible for tissue fluid homeostasis, lipid absorption, and mediation of immune responses. It initiates as a network of blind-ended capillaries throughout most organs of the body. These capillaries have discontinuous tight junctions in order to take up free fluid and solutes from the interstitium or lipids from the intestinal tract. Antigen-presenting cells, as well as antigen itself, enter the lymphatic system at the level of the capillaries before being conducted into collecting lymphatic vessels (LVs). Concurrent with an increasing diameter, collecting LVs are covered by a thin layer

of smooth muscle cells that are required for the intrinsic contractility of lymphatic collectors (Gashev, 2008; Zawieja et al., 1993). Together with intraluminal valves, their active contractions ensure unidirectional lymph flow against an increasing pressure gradient. Collecting LVs conduct lymph through one or more lymph nodes until it ultimately reaches the bloodstream at the subclavian veins.

The mechanisms regulating collecting LV contractility are not entirely understood. Generally, there are both extrinsic and intrinsic factors that influence lymph flow. Extrinsic factors include the passive movement of lymph by contractions of striated muscles (e.g., during walking), as well as by nearby arteries, which through vasomotion can conduct their pulsation to LVs (Gashev, 2008; Zawieja et al., 1993). With regard to intrinsic mechanisms, mechanical parameters such as shear stress and intraluminal pressure gradients within the LVs exert important influences on smooth muscle cell behavior (reviewed in Scallan et al., 2016). Vasoregulatory factors, including nitric oxide (NO), also have a strong influence on the contractility. NO is an effective vasodilator and has negative inotropic effects on LVs (Scallan and Davis, 2013). The release of NO is induced by high shear stress within the vessel (Kornuta et al., 2015; Kunert et al., 2015) or by a variety of signaling molecules such as vascular endothelial growth factor A (VEGF-A) (Lahdenranta et al., 2009). Surprisingly, the potential regulation of LV contractility by the autonomic nervous system has not received much attention thus far.

Nerves in close proximity to LVs have been described in humans (D'Andrea et al., 2013, 2015) and large animals, including cows and dogs (Ohhashi et al., 1982; Todd and Bernard, 1973), and *ex vivo* studies have reported responses of the contraction pattern of isolated LVs to several neurotransmitters (McHale et al., 1980, 1990; Ohhashi and Azuma, 1986). However, the results of these *ex vivo* studies are somewhat contradictory. While previous studies consistently found activation of lymphatic contractility by the α -adrenergic nervous system, some attributed these effects solely to the activation of α_1 receptors (Benoit, 1997), whereas others reported that only α_2 receptors, but not α_1 receptors, are needed (Hashimoto et al., 1994). Similarly, conflicting results have been reported regarding the role of the



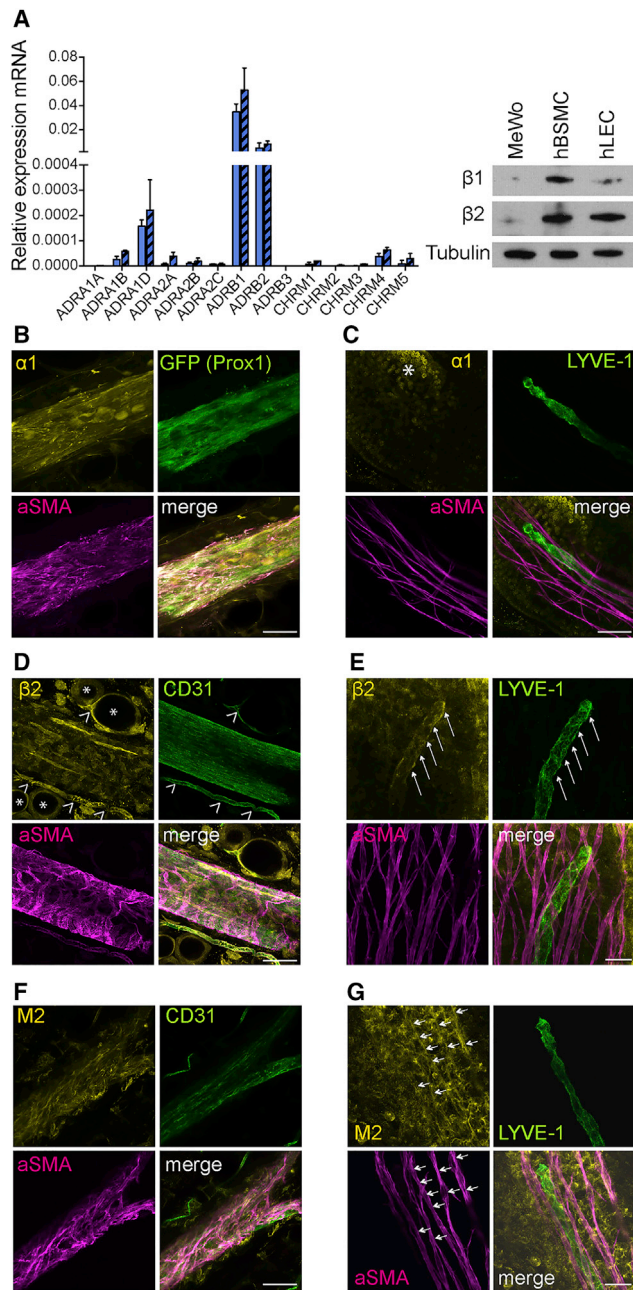


Figure 1. Neurotransmitter Receptors Are Expressed on LVs in an Organ-Specific Manner

(A) qPCR-based detection of adrenergic β_1 and β_2 receptor (*ADRB1* and *ADRB2*) mRNA in two primary human LEC lines (shown as blue and blue-striped columns). Expression was confirmed at the protein level using western blots with human bronchial smooth muscle cells (hBSMCs) as a positive control and the tumor cell line MeWo as a negative control.

(B, D, and F) Whole-mount staining of flank collector showing α_1 (B), β_2 (D), and M2 (F) receptors expressed on CD31⁺ LECs and α SMA⁺ smooth muscle cells. Asterisks indicate β_2 -positive CD31⁺ blood vessels, and open arrowheads indicate β_2 -positive adipocytes.

(C, E, and G) LYVE-1⁺ lacteals and their associated α SMA⁺ cells do not express α_1 receptors (C), but the lacteals express β_2 receptors (E, arrows), and the associated smooth muscle cells express M2 receptors (G, arrows). The α_1

muscarinic nervous system, since its agonist acetylcholine (ACh) induced LV relaxation (Ferguson, 1992; Hashimoto et al., 1994; Ohhashi and Takahashi, 1991) as well as contraction (Ohhashi et al., 1978). Recently, *ex vivo* studies using human tissue found a strong induction of LV contracting activity by the adrenergic agonist norepinephrine (NE) and an inhibitory effect of the muscarinic antagonist atropine (Telinius et al., 2010, 2014). Overall, most of the current knowledge about the interaction between the autonomic nervous system and LV contractility stems from *ex vivo* studies, while *in vivo* studies are still lacking (Choe et al., 2015; Ono et al., 2000). Importantly, virtually nothing is known about potentially different responses of LVs to neurotransmitters in pathological conditions such as chronic inflammation and cancer, where LVs play major pathogenetic roles (Christiansen and Detmar, 2011; Stacker et al., 2014).

In this study, we investigated in detail the role of the autonomic nervous system in the regulation of LV function under physiological and pathological conditions. To this end, we first studied the presence of different subtypes of nerves on murine LVs and analyzed the expression profile of neurotransmitter receptors on lymphatic endothelial cells (LECs) *in vitro* and *in situ*. These studies revealed organ-specific expression patterns that might explain previously reported diverging effects of neurotransmitters in different organs. We then applied quantitative *in vivo* near-infrared imaging techniques to directly investigate the effects of a broad spectrum of agonists and antagonists of the autonomic nervous system on the contraction frequency and amplitude of collecting LVs under physiological conditions. These investigations were accompanied by live Ca²⁺ imaging studies *in vitro* and *in vivo*. Finally, we investigated the effects of selected neurotransmitters on tumor-draining LVs. Our results reveal that under physiological conditions, contractions of collecting LVs are activated via α -adrenergic and muscarinic receptors, while they are inhibited via activation of β_2 -adrenergic receptors. Importantly, we found that tumor-draining LVs exhibited more active contractility and that this contraction activity can be inhibited by distinct agonists and antagonists of the autonomic nervous system.

RESULTS

Neurotransmitter Receptors Are Expressed on LVs in an Organ-Specific Manner

We first investigated the mRNA expression levels of all known adrenergic and muscarinic neurotransmitter receptors in five different primary human dermal LEC lines. LECs specifically expressed β_1 - and β_2 -adrenergic receptors (Figure 1A shows two cell lines, and Figure S1A the additional three lines). These findings were confirmed at the protein level by western blot (Figure 1A), with human bronchial smooth muscle cells (hBSMCs) serving as a positive control and the tumor cell line MeWo as a negative control. In a next step, we investigated the neurotransmitter receptors expressed in mouse tissue. Whole-mount

receptor expressing enterocytes (Baglole et al., 2006) are clearly visible (asterisk in C). Images were obtained using confocal microscopy.

Scale bars represent 50 μ m (B, D, and F) and 25 μ m (C, E, and G). Data in (A) are presented as mean \pm SD. See also Figure S1.

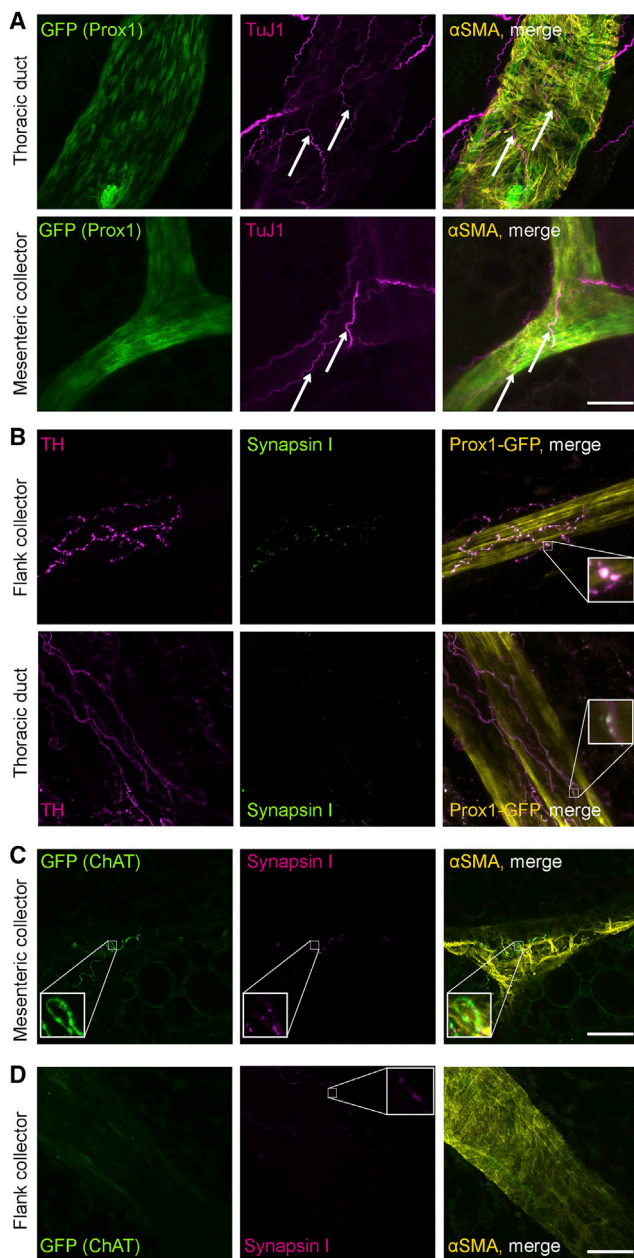


Figure 2. Nerves Are Closely Associated with LVs

(A) Lymphatic collectors of *Prox1-GFP* mice are closely associated with pan-neuronal marker TuJ1⁺ nerves (arrows).

(B) Tyrosine hydroxylase (TH)⁺ sympathetic nerves run along lymphatic collectors of *Prox1-GFP* mice and form synapses (synapsin I, zoom-in).

(C) Choline acetyltransferase (ChAT) GFP⁺ parasympathetic nerves, which form synapses (synapsin I), accompany mesenteric collectors (zoom-in).

(D) Collectors other than mesenteric collectors show no ChAT⁺ innervation. Synapsin I staining shows the presence of other types of nerves (zoom-in). Images were obtained using confocal microscopy.

Scale bars, 50 μ m. See also Figure S2.

staining of the flank collecting LVs and lacteal villi in the small intestine was performed. Adrenergic α_1 receptors were present on CD31⁺ LECs as well as on the α SMA⁺ smooth muscle cells of the

flank collector (Figure 1B). In contrast to the collecting LVs, neither the intestinal LYVE-1⁺ lacteals nor their associated smooth muscle cells (α SMA⁺) demonstrated expression of α_1 receptors (Figure 1C). Interestingly, the mesenteric collecting LVs lacked expression of α_1 receptors (Figure S1F). Whole-mount staining revealed no detectable expression of β_1 receptors on either type of LVs (Figures S1B and S1C), whereas the β_2 receptors were expressed on LECs of all vessel types, lacteals, the flank, and mesenteric collectors (Figures 1D, 1E, and S1G). β_2 -Receptor expression was also detected on smooth muscle cells of the flank and mesenteric collecting LVs, but not on the smooth muscle cells surrounding the lacteal.

Even though it was not detected at the RNA level in cultured human LECs, we next performed whole-mount staining for the muscarinic receptor M₂. We found expression of M₂ receptors by smooth muscle cells of the flank and mesenteric collector and of the lacteal villi, whereas the LECs themselves only showed expression of M₂ receptors in the collecting vessels (Figures 1F, 1G, and S1H). As a second method, flow-cytometry studies on freshly isolated flank collector LECs were performed (Figures S1I and S1J). Thereby, we were able to confirm the expression of α_1 and β_2 receptors. Additionally, but to a lower extent than β_2 receptors, we also detected positive expression of β_1 receptors. Due to the lack of a suitable antibody, we were unable to assess M₂ receptor expression using this approach. Taken together, these findings reveal that LVs specifically express receptors in an organ-site-dependent manner that may enable them to respond to neurotransmitters released from the autonomic nervous system.

Nerves Are Closely Associated with LVs

In order to exert neural effects on LVs, the presence of synapses formed by nerves is required, as these represent the source of neurotransmitters. Whole-mount staining of collecting LVs from different organ sites of *Prox1-GFP* mice for the pan-neuronal marker TuJ1 detected nerves running along LVs (Figure 2A). Tyrosine hydroxylase (TH)-positive sympathetic nerves were detected, forming synapsin-I-positive synapses on lymphatic collectors (Figure 2B). Parasympathetic nerves are characterized by their expression of choline acetyltransferase (ChAT). We therefore performed whole-mount staining of *CHAT-GFP* mice in combination with synapsin I staining to detect synapses. Mesenteric lymphatic collectors demonstrated innervation with parasympathetic nerves (Figure 2C, zoom-in), whereas no ChAT⁺ neural structures were detected on collectors at other organ sites, such as the flank (Figure 2D).

Whole-mount staining showed a close proximity of nerves (TuJ1⁺), distinguishable as either sympathetic (TH⁺) or parasympathetic (ChAT⁺), with intestinal lacteals (Figures S2A–S2C). Synapsin I staining revealed that the nerves adjacent to lacteals also formed synapses (Figures S2D and S2E).

Overall, these results demonstrate that autonomic nerves of different subtypes are present near LVs, indicating the possibility of a direct influence on LV function.

LECs Are Direct Targets of Neurotransmitters

To evaluate if LECs might demonstrate downstream signaling in response to neurotransmitters, we next investigated their Ca²⁺

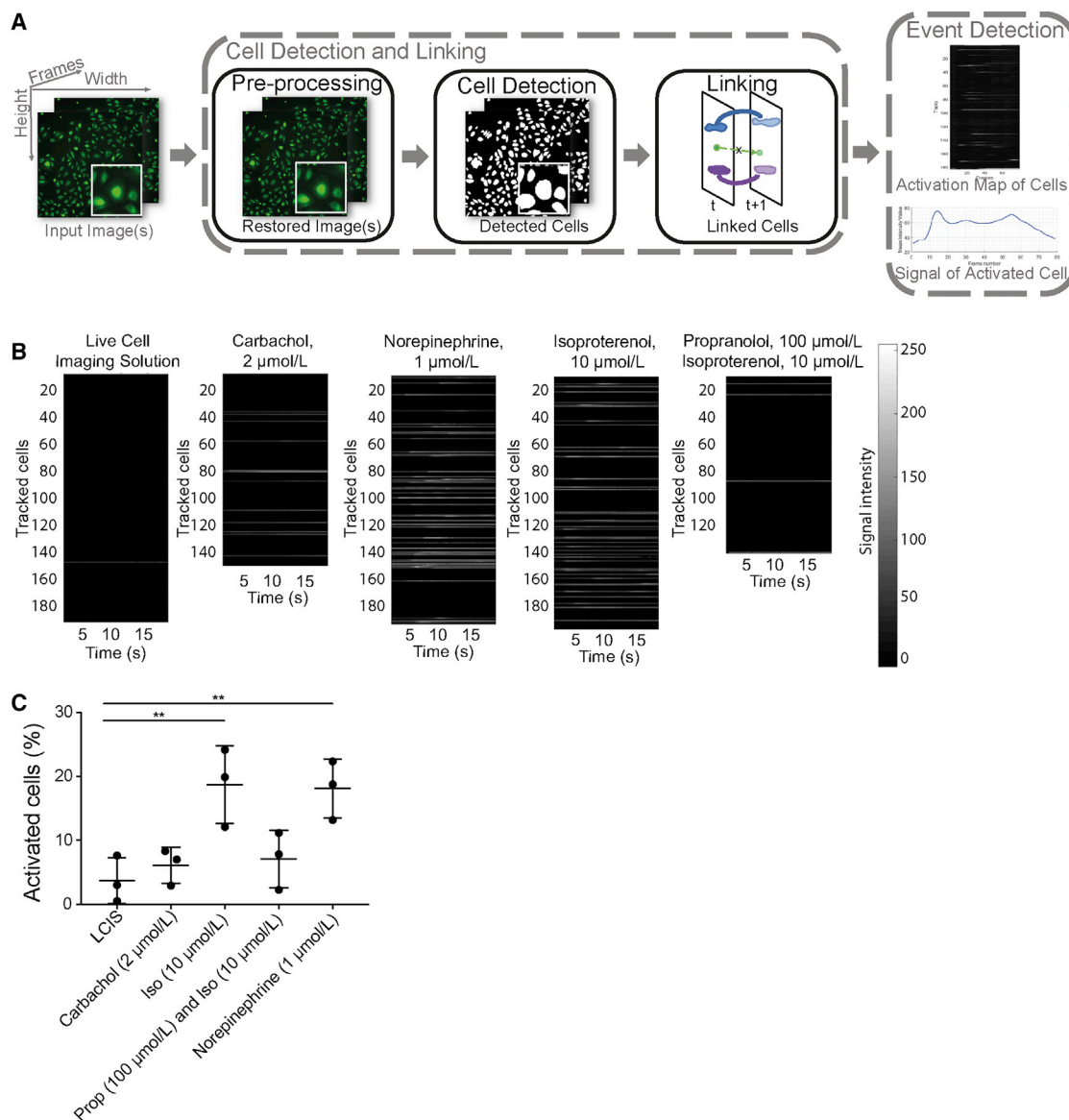


Figure 3. LECs Are Direct Targets of Neurotransmitters *In Vitro*

(A) Visualization of the algorithm made for Ca^{2+} quantification. After enhancing the quality of each frame (image pre-processing), each cell is detected separately and linked between consecutive frames in order to track it over the whole duration of the video (Cell Detection and Linking). For each tracked cell (Track), the fluorescence intensity over time is computed. If the signal changes notoriously over time, an activation event is recorded (Event Detection). A detailed description can be found in [STAR Methods](#).

(B) Quantification of *in vitro* Ca^{2+} imaging of human primary LECs using Fluo8H in response to different neural agonists. Each horizontal line of the bar represents a tracked cell; only activated cells are shown in gray. Fluo8H intensity is shown in grayscale, proportional to intracellular Ca^{2+} .

(C) Quantification of the percentage of cells reacting with an increase of Ca^{2+} upon treatment with different agonists. LCIS, live cell imaging solution; Iso, isoproterenol; Prop, propranolol. ** $p \leq 0.01$. Data in are presented as mean \pm SD; n = average per well of cells. Individual cells can be seen [Figure S3](#). See also [Figure S3](#).

response *in vitro* as well as *in vivo*. This approach was chosen because activation of most muscarinic receptors and some adrenergic receptors leads to an increase of intracellular calcium as a second messenger. *In vitro* imaging of human LECs using Fluo8H was quantified with a self-written algorithm that is explained in [STAR Methods](#) and [Figure 3A](#). We found that intracellular Ca^{2+} was increased upon treatment with NE (1 $\mu\text{mol/L}$) or

isoproterenol (10 $\mu\text{mol/L}$) ([Figures 3B](#) and [S3](#)). To evaluate the specificity of this experiment, pretreatment of cells with an antagonist was performed. Propranolol (100 $\mu\text{mol/L}$) strongly inhibited the increase of intracellular Ca^{2+} induced by isoproterenol ([Figure 3B](#)). The quantification of the percentage of activated cells was in line with the visual quantification of Ca^{2+} activation ([Figure 3C](#)).

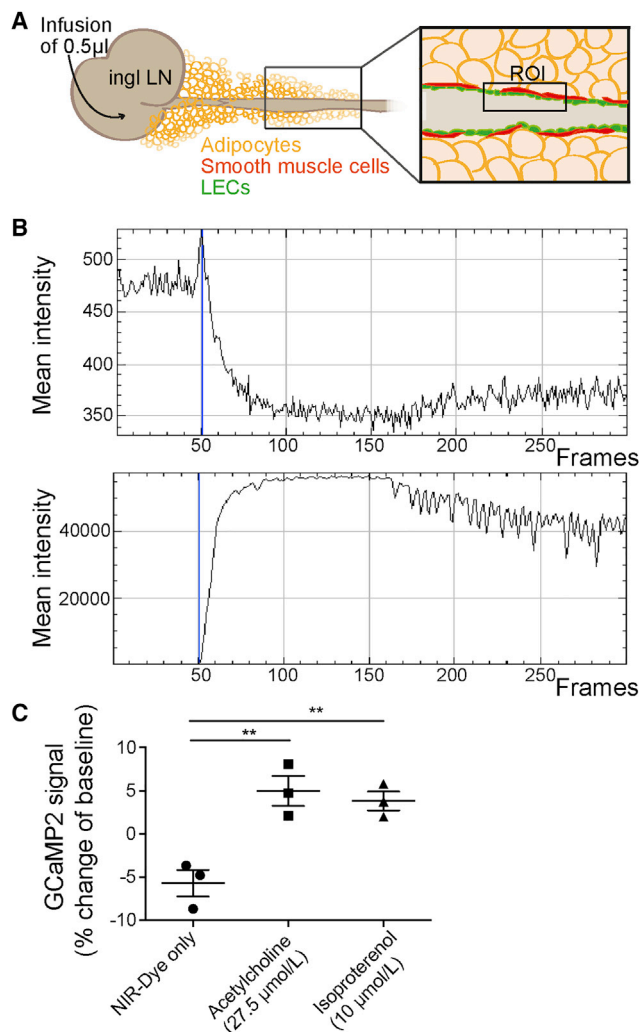


Figure 4. LECs Are Direct Targets of Neurotransmitters In Vivo
 (A) Schematic overview of *in vivo* Ca^{2+} imaging using *Cx40* *GCaMP2* mice. The flank collector was used for imaging. Ingl LN, inguinal lymph node; ROI, region of interest.
 (B) Representative example of *GCaMP2* intensity (top) measurement while a mixture of pegylated NIR dye and acetylcholine was infused into the inguinal lymph node. At time of arrival of the dye (bottom), there is a visible peak of the *GCaMP2* expression (frame 50, blue line).
 (C) The *GCaMP2* signal intensity at time point of arrival of the dye in the imaged section was normalized to the signal intensity before arrival of the dye. Differences are shown as percent change of the baseline signal intensity. ** $p < 0.01$. Data are presented as mean \pm SD; $n = 3$ mice.

For *in vivo* imaging, *Cx40* *GCaMP2* transgenic mice were used (Tallini et al., 2007). These mice express the calcium indicator *GCaMP2* under control of the connexin 40 promoter and provide a dynamic readout of GFP signal intensity that is proportional to the intracellular calcium concentration. Connexin 40 is expressed in arterial endothelial cells and to a lesser degree in LECs. In these mice, we infused 0.5 μ L of 10 μ mol/L 20 kDa PEG-IRDye680 (P20D680) near-infrared (NIR) tracer (Proulx et al., 2013) either alone or mixed with adrenergic or muscarinic agonists into the inguinal lymph node (Figure 4A) and evaluated a

region of interest over the efferent collecting LV. The tracer was used to define the time point when the infused agonist arrives in the region of interest. Ca^{2+} fluxes in the LECs were analyzed by measuring changes in the GFP intensity before arrival of the tracer compared to the time point of arrival (Figure 4B). When tracer alone was infused, the GFP intensity dropped due to the volume-induced dilation of the collecting vessel (Figure 4C). When the muscarinic agonist ACh (27.5 μ mol/L) was infused with the tracer, a clear transient spike in GFP intensity was observed (Figure 4C). The same was observed for the β adrenergic agonist isoproterenol (10 μ mol/L).

Together, these results indicate that LECs might represent direct targets of neurotransmitters.

The Autonomic Nervous System Regulates Lymphatic Contractility In Vivo

To investigate the regulation of LV contractility by the autonomic nervous system *in vivo*, we used a previously described method (Chong et al., 2016). In brief, after the infusion of 0.5 μ L of the P20D680 NIR dye into the inguinal lymph node, the flank collector becomes visible and contractile activity can be imaged (Figure 5A). Over a period of 8 min, the fluorescent intensity of the tracer in the vessel is imaged, which corresponds to diameter changes of the vessel (Figure 5B) and enables quantification of the contractility using a custom-made algorithm (Chong et al., 2016). After the first 2 min, a topical treatment was applied. We quantified contraction frequency, amplitude, and pumping score, which is the product of frequency and amplitude, after treatment with a range of muscarinic, α -adrenergic, and β -adrenergic agonists and antagonists.

Comparing the post-treatment period (minutes 4–7) with the pretreatment period (first 2 min), potent effects of several agonist and antagonists were found (Figures 5C–5E, S5C, and S5D). There was a strong increase of contractile activity upon treatment with the muscarinic agonist carbachol (2 μ mol/L; amplitude 212.62% of pre-value; pumping score 317.67% of pre-value). Conversely, antagonization of the muscarinic system by atropine (100 nmol/L) reduced the contraction activity significantly (frequency 57.63% of pre-value; pumping score 44.26% of pre-value). For the adrenergic system, we detected reduced contractility upon treatment with the unspecific β -adrenergic agonist isoproterenol (10 μ mol/L; frequency 45.1% of pre-value). To determine if this effect was mediated by β_1 or β_2 receptors, the specific β_1 agonist denopamine and the specific β_2 agonist salbutamol (both at 1 μ mol/L) were studied. Only salbutamol induced an inhibitory effect (frequency 65.29% of pre-value), indicating that the effect is predominantly mediated by β_2 receptors. Treatment with agonists at two additional concentrations (0.1 and 10 μ M) also showed no significant effect of the β_1 agonist, while the higher concentration of the β_2 agonist showed a strong trend of reduced frequency (62.88% of pre-value) (Figure S5B). The unspecific β adrenergic antagonist propranolol (10 μ mol/L) exhibited a trend to activate the contraction activity (frequency 115.07% of pre-value, amplitude 171.55% of pre-value, pumping score 197.86% of pre-value). To ensure specificity of the β_2 agonist, competitive blocking studies were performed. After the exposed vessels were treated at $t = 2$ min,

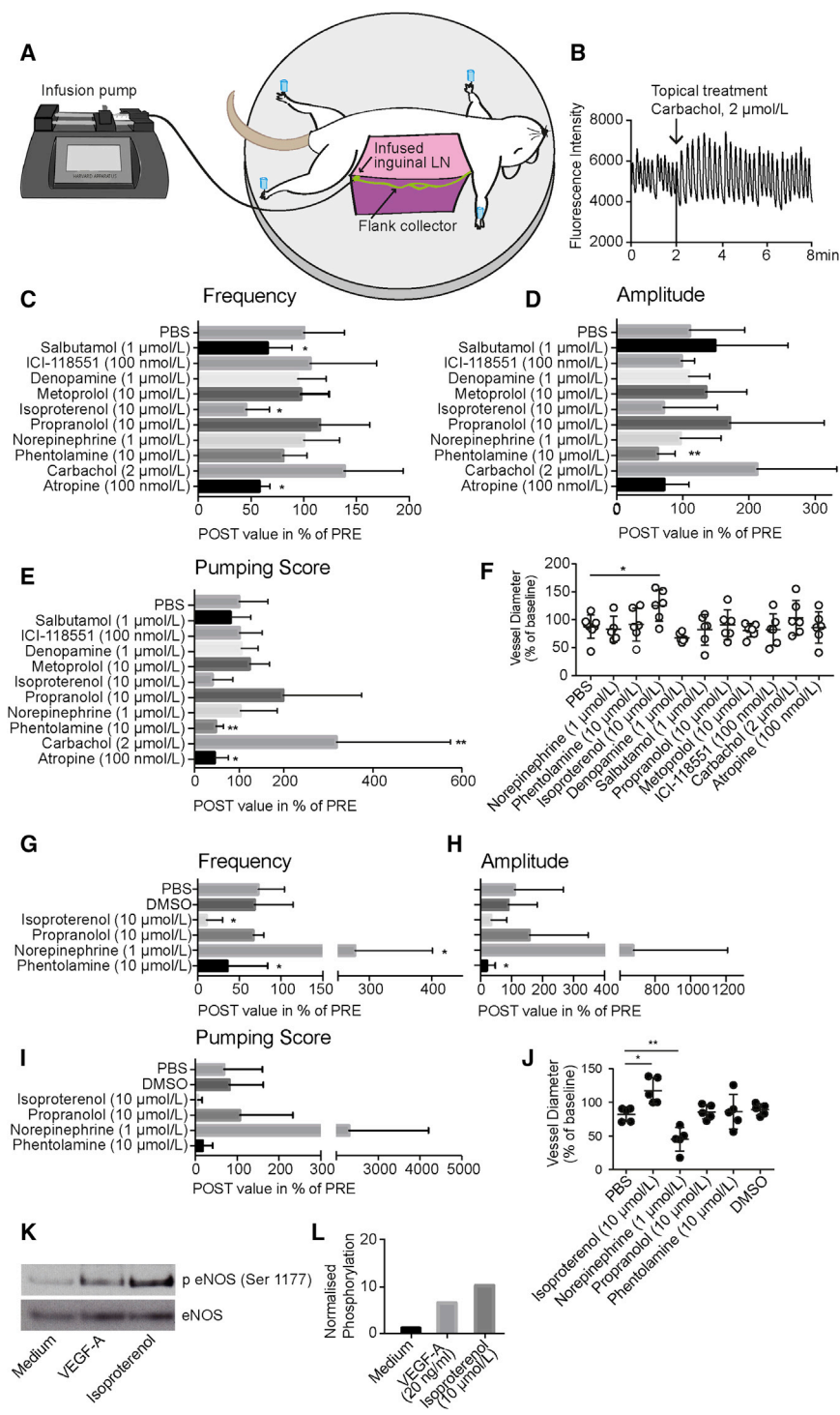


Figure 5. The Autonomic Nervous System Regulates Lymphatic Contractility In Vivo

(A) Schematic overview of *in vivo* imaging of flank collector contractility.

(B) Representative example of measured NIR-dye intensity in the flank collector over time; topical treatment was done at time point 2 min.

(C–E) Effects of topical treatment with neural agonists and antagonists on lymphatic contraction frequency (C), amplitude (D), and pumping score (E). Shown are the post-treatment values as a percentage of pretreatment values for mice under injection anesthesia (ketamine, xylazine, and acepromazine).

(G–I) Same is shown for mice under isoflurane: changes in frequency (G), amplitude (H) and pumping score (I).

(F and J) Quantification of the tonic contraction of the flank collector upon treatment with neural agonists and antagonists under injection anesthesia (F; ketamine, xylazine, acepromazine) and isoflurane (J).

(K and L) Isoproterenol-mediated phosphorylation of eNOS, as assessed (K) and quantified (L) by western blot. VEGF-A was used as positive control. The western blot shown is a representative example of at least three independent experiments.

* $p < 0.05$, ** $p \leq 0.01$. Data are presented as mean \pm SD. (C–F) $n = 8$ vessels for PBS, and $n = 6$ vessels for all other treatments. (G–J) $n = 4$ vessels per treatment. (K and L) $n = 1$. See also Figure S5.

The α -adrenergic system has been reported to induce contractility in isolated LVs (Ono et al., 2000; Telinius et al., 2014). In line with these results, we found a reduced contraction activity upon antagonization of the α adrenergic system using phentolamine (10 μ mol/L; amplitude 62.42% of pre-value; pumping score 46.79% of pre-value; Figures 5D and 5E). Phentolamine was dissolved in DMSO, reaching a final DMSO concentration of 0.01%. In previous studies, we confirmed that this concentration of DMSO had no effect on the contraction activity (Chong et al., 2016). Surprisingly, we could not detect any effect of the α -adrenergic (and, with much lower affinity, β -adrenergic) agonist NE. This lack of response was most probably due to the usage of the injection anesthesia (ketamine, xylazine, and acepromazine) that contained α agonists itself. To investigate this hypothesis, we repeated the adrenergic treatments with mice under isoflurane anesthesia (Figures 5G–5I and S6). We confirmed the inhibiting effect of the α antagonist phentolamine (10 μ mol/L; amplitude 19.87% of pre-value), while the DMSO control showed no significant effect. More importantly, we found a strong activating

either PBS or the β_2 antagonist were added at $t = 4$ min, and the vessels' post-treatment was assessed at $t = 6$ –9 min. While PBS treatment had no effect on the reduction of contractility, treatment with the β_2 antagonist neutralized the effect of the initial β_2 agonist, resulting in no change in contractility (Figure S5A).

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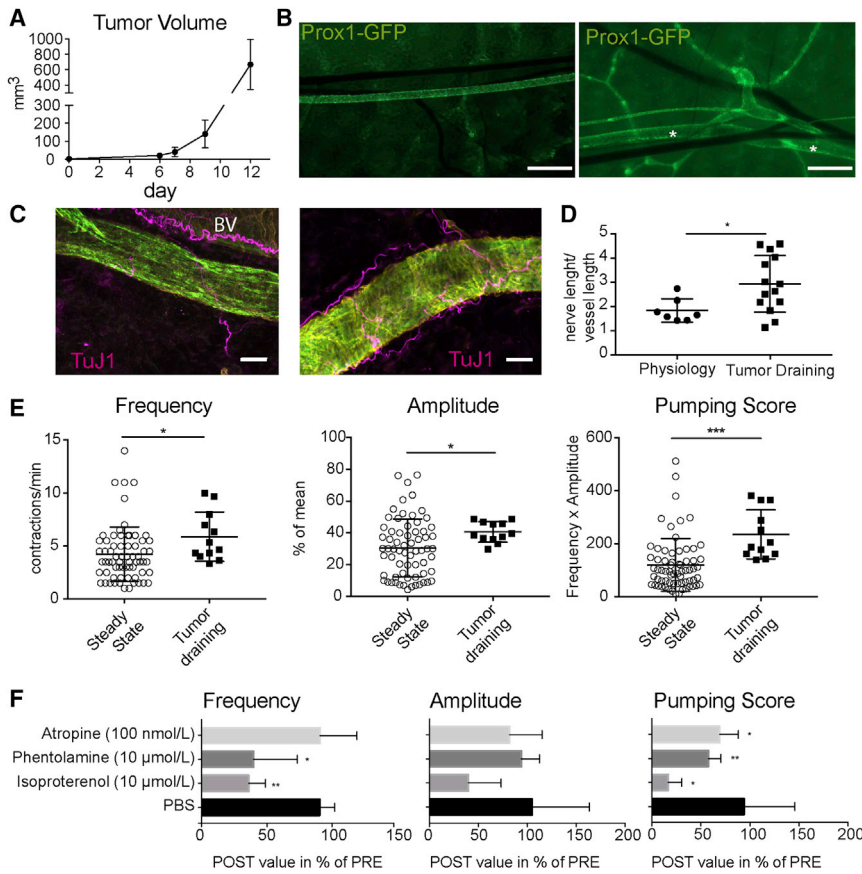


Figure 6. Tumor-Draining Lymphatic Collectors Contract More Actively

(A) Growth of intradermally injected B16F10 tumors was measured over time using a caliper ($n = 7$ mice).

(B) Overview picture of the flank area in a naive (left) and a tumor-bearing *Prox1-GFP* mouse (right). In the vicinity of the flank collector (asterisk), additional lymphatic collectors were formed draining the tumor.

(C) Compared to physiological LVs (left), tumor-draining vessels show denser innervation, visualized with the pan-neural marker TuJ1 (*Prox1-GFP* staining, green; α SMA, yellow).

(D) Quantification of innervation density (nerve length/vessel length).

(E) Tumor-draining LVs show higher baseline activity than the respective vessels in non-tumor-bearing mice with regard to frequency, amplitude, and pumping score.

(F) Contractility of tumor-draining vessels can be inhibited by atropine, phentolamine, and isoproterenol.

* $p < 0.05$, ** $p \leq 0.01$, and *** $p \leq 0.001$. Scale bars represent 500 μm (B) and 25 μm (C). Data represent mean \pm SD, and dots in (D) and (E) represent vessels. (F) $n = 4$ vessels per treatment. BV, blood vessel. See also Figure S6.

effect of NE (1 $\mu\text{mol/L}$; frequency 277.29% of pre-value), indicating that the lack of effect in the previous experiments was due to the choice of anesthesia. To investigate whether the results observed after treatment with other mediators might also have been affected by the injection anesthesia, we next treated collecting LVs in mice anesthetized with isoflurane with β adrenergic compounds. Isoproterenol (10 $\mu\text{mol/L}$) induced comparable effects (frequency 10.32% of pre-value) and propranolol (10 $\mu\text{mol/L}$) had only minor effects on the contractility (frequency 66.63% of pre-value) (Figures 5G–5I).

Overall, these *in vivo* experiments identified a strong impact of neurotransmitters on the contraction activity of lymphatic collectors in mice.

β -Adrenergic Stimulation Regulates LV Tone by eNOS Phosphorylation

Besides the effects on phasic contractions, we next investigated whether treatment with neurotransmitters might also mediate changes in vessel tone, since previous studies indicated that NE increases the vessel tone of lymphatic collectors from various organ sites by inducing vasoconstriction (Dobbins, 1992; Ono et al., 2000; Takahashi et al., 1990). While mice under injection anesthesia did not show any effect of NE on the vessel tone (Figure 5F), in line with the lack of any response in phasic contractions, the vessel tone was increased in mice under isoflurane anesthesia (Figure 5J). We observed a decrease in vessel

tone (vasodilation) upon treatment with isoproterenol, independent of the anesthesia regimen used. As a potential mediator of vasodilation, we next investigated the phosphorylation state of eNOS in LECs upon treatment with β -adrenergic agonists by western blot and found an increased phosphorylation after treatment with isoproterenol (Figures 5K and 5L). Thus, treatment with isoproterenol may lead to increased NO production in LECs. The released NO acts on the nearby smooth muscle cells, induces their relaxation, and, therefore, leads to a dilation of the collecting LV.

Tumor-Draining Lymphatic Collectors Contract More Actively

B16F10 melanomas were grown in the flank skin of *Prox-1 GFP* mice (Figure 6A). When tumors reached 12 mm in one dimension, tumor-draining LVs were imaged *in vivo*. As reported by many studies (Padera et al., 2002; Skobe et al., 2001; Stacker et al., 2014), we found that several LVs were expanded in the vicinity of the flank collector, demonstrating the direct activating effect of the tumor on the lymphatic vasculature (Figure 6B). TuJ1 staining of the affected vessels showed a denser innervation (Figures 6C and 6D). In line with this finding, the tumor-draining flank collectors had a higher baseline contraction activity (Figure 6E). Based on the results obtained under physiological conditions, we next investigated whether treatment with distinct neurotransmitters might inhibit the contractility of tumor-draining LVs. Indeed, atropine, phentolamine, and, most potently, isoproterenol reduced the pumping score significantly (Figures 6F and S6).

DISCUSSION

In this study, we provide the first comprehensive *in vivo* characterization of the regulation of LV contractility by agonists and antagonists of the autonomic nervous system. Our findings reveal that both smooth muscle cells and endothelial cells of LVs express distinct neurotransmitter receptors and that nerves and synapses are in contact with these vessels. We identified specific functions of the sympathetic and the parasympathetic nervous system with regard to LV contraction and confirmed the distinct activation of signaling in LECs by Ca^{2+} imaging *in vitro* and *in vivo*. The studies of tumor-draining LVs revealed their increased innervation and contraction activity and also identified neurotransmitters that may inhibit the activity of tumor-draining LVs.

In our investigation of the expression of neurotransmitter receptors in LECs, we found that cultured human LECs only express β -adrenergic receptors, whereas, in addition, both α_1 -adrenergic and M_2 -muscarinic receptors were found on murine collecting LVs by whole-mount staining and flow-cytometry analysis. The lack of expression of several of these receptors *in vitro* might indicate a change of the expression profile of LECs under the *in vitro* culture conditions, in agreement with previous studies (Amatschek et al., 2007; Wick et al., 2007). In addition, these differences might also represent vessel-type-specific expression patterns, since the cultured human LECs were isolated from the skin and might be more representative of LECs originating from capillaries and not from collecting LVs. Consistent with this concept, murine lacteal LECs, representing cells of specialized intestinal lymphatic capillaries, also only express β -adrenergic receptors. However, species-associated differences between humans and mice cannot be ruled out as well. Confirmation for the differently expressed receptors in LECs *in vitro* compared to the *in vivo* situation was obtained by the Ca^{2+} imaging studies, since cultured cells only responded to β agonists, whereas there was also a response to the muscarinic agonist ACh *in vivo*.

Our finding of an organ-specific expression pattern of neurotransmitter receptors may provide an explanation for the previously reported discrepancies of NE effects on LV function. While contractility was increased in *ex vivo* preparations of peripheral collecting LVs from different species (including human) after exposure to NE (Dobbins, 1992; Ono et al., 2000; Takahashi et al., 1990; Telinius et al., 2014), in agreement with our *in vivo* studies in mice, a recent study reported a diminished contraction activity of intestinal lacteals after NE treatment (Choe et al., 2015). Our findings indicate that the opposing effects of the same adrenergic agonist can be explained by the absence of α -adrenergic receptors on the lacteals. Thus, NE (being an α -adrenergic agonist and, with much lower affinity, also a β -adrenergic agonist) likely acts as a β agonist in lacteals, leading to reduced contractility, whereas it promotes contractions in lymphatic collectors via the α -adrenergic receptors. This organ-specific response to NE makes sense physiologically, since under systemic stress conditions, enhanced return of peripheral lymph fluid together with reduced energy expense for the gastrointestinal lipid transport might be advantageous.

A major conclusion from our studies is that muscarinic and α -adrenergic agonists activate the contractility of collecting LVs, whereas β_2 -adrenergic agonists reduce the contraction frequency. The activating effect mediated by NE is in agreement with previous *ex vivo* studies (Hashimoto et al., 1994; Igarashi et al., 1998; McHale, 1992; McHale et al., 1980; Telinius et al., 2014). However, whereas some of these studies indicated that this effect is mediated via α_2 receptors (Hashimoto et al., 1994), others suggested that both α_1 and α_2 receptors are involved (Igarashi et al., 1998). Our findings that there are profound differences in the expression patterns of neurotransmitter receptors on different types of LVs indicate that some of the reported differences might relate to organ-specific responses to neurotransmitters. In support of this concept, LVs isolated from the liver and lungs of pigs showed decreased contraction and were dilated upon ACh treatment (Ferguson, 1992; Hashimoto et al., 1994), similarly to isolated thoracic ducts of dogs (Ohhashi and Takahashi, 1991). By contrast, in rat mesenteric vessels (Fang et al., 2007), operation-induced vagotomy resulted in a reduced contraction rate, indicating an activating role of the parasympathetic nervous system with its main neurotransmitter, ACh, in line with the contraction-inducing effects of ACh in bovine mesenteric LVs (Ohhashi et al., 1978). Another possible explanation for these opposing effects of ACh might be the ratio of muscarinic receptors expressed on LECs versus smooth muscle cells. If muscarinic receptors are expressed predominantly on smooth muscle cells, then their activation likely induces LV contraction, whereas binding of ACh on muscarinic receptors on LECs induces NO release (Scallan and Davis, 2013), resulting in vessel dilation and reduced contractility. Thus, a detailed characterization of the neurotransmitter receptor expression profiles of human LVs from different organs, as well as from model organisms such as the mouse, is needed to better understand the organ- and species-specific effects of the autonomic nervous system.

Our data indicate that the choice of anesthesia is of great importance for the design and interpretation of functional studies of LV contractility. We were unable to detect the activating effects of the α -agonist NE when using an injection anesthesia regimen that contained the α -agonist xylazine, thus masking the effects of NE on LV contractility. By contrast, NE induced an increased LV contraction frequency under isoflurane anesthesia. However, this anesthesia regimen turned out to be associated with a general depression of LV contractions. Thus, a thorough investigation of different anesthesia regimens for functional *in vivo* studies of LVs would be helpful to ensure a better cross comparison of published work.

The results of our *in vivo* studies are in agreement with the improvement of secondary lymphedema after therapeutic inhibition of the sympathetic nervous system in patients after breast cancer surgery (E. Choi et al., 2015) and gynecologic-cancer-related surgery (Woo et al., 2013). In these clinical settings, the treatment blocks the complete sympathetic nervous system locally. Thus, one might speculate that the parasympathetic nervous system, which is left intact after the treatment, promotes LV contractions, resulting in improvement of the lymphedema.

One should keep in mind that the autonomic nervous system influences both lymphatic and blood vessels, indicating that

therapies aimed at modulating blood vessel functions might also impact the function of the lymphatic vascular system. As an example, the long-term usage of β blockers for hypertension might negatively influence the balanced system of physiological lymphatic contractions. Similarly, while β_2 agonists are successfully used to induce bronchodilation in asthma patients, they might reduce the contractility of lymphatics of the lung and increase NO release by LECs, potentially leading to a reduced fluid clearance of the lung tissue.

To our knowledge, our studies identified for the first time that the expanded tumor-draining LVs have a higher innervation density, which might contribute to the observed increase in contractility of the preexisting and newly formed LVs. The increased contractility of tumor-draining LVs is in agreement with a previous study in mice where increased smooth muscle cell coverage was also detected (Gogineni et al., 2013). Together, these findings reveal that tumors not only induce LEC proliferation and lymphangiogenesis but also profoundly change the anatomical structure and function of tumor-draining LVs. Our findings that specific agonists and antagonists of the autonomic nervous system reduce the contractility of tumor-draining LVs might be of potential relevance for clinical cancer therapy. In the future, targeted long-term treatment of tumor-draining LVs would be needed to investigate whether the inhibition of lymphatic contractility ultimately results in reduced tumor growth and metastasis. At the same time, these studies could also address possible negative effects of an increased interstitial pressure induced by such long-term treatments, including potentially reduced tumor tissue penetration of therapeutic compounds.

STAR★METHODS

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SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at <https://doi.org/10.1016/j.celrep.2019.05.050>.

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AUTHOR CONTRIBUTIONS

Conceptualization, S.B.B. and M.D.; Software, J.M., F.S., and S.F.N.; Investigation, S.B.B., D.G., M.S., S.T.P., and C.T.; Resources, M.D.; Writing – Original draft, S.B.B.; Writing – Review & Editing, all authors; Supervision, M.D. and S.F.N.; Funding Acquisition, M.D.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
alpha-1 adrenergic receptor	Santa Cruz	cat# sc-1477; RRID:AB_630854
beta-1 adrenergic receptor	abcam	cat# ab3442; RRID:AB_10890808
beta-2 adrenergic receptor	abcam	cat# ab13989; RRID:AB_300816
muscarinic acetylcholine receptor 2	abcam	cat# ab109226; RRID:AB_10858602
muscarinic acetylcholine receptor 3	Santa Cruz	cat# sc-9108; RRID:AB_2291779
Synapsin-1	Invitrogen	cat# 51-5200; RRID:AB_87668
Tyrosine Hydroxylase	R&D	cat# AF7566
Tyrosine Hydroxylase	Invitrogen	cat# P21962; RRID:AB_2539844
III beta Tubulin	BioLegend	cat# 801201; RRID:AB_2313773
GFP	Aves	cat# GFP-1010; RRID:AB_2307313
CD31	BD Bioscience	cat# 550274; RRID:AB_393571
alpha smooth muscle actin	Sigma-Aldrich	cat# C6198; RRID:AB_476856
LYVE-1	AngioBio	cat# 11-034
LYVE-1	ReliaTech	cat# 103-M130
eNOS	ThermoFisher	cat# PA1-037; RRID:AB_325774
Phospho-eNOS (Ser1177)	CellSignaling	cat# 9571; RRID:AB_329837
Tubulin	Sigma-Aldrich	cat# T9026V; RRID:AB_477593
goat Alexa 594	Invitrogen	cat# A-11058; RRID:AB_2534105
rabbit Alexa 488	Invitrogen	cat# A-21206; RRID:AB_2535792
rabbit Alexa 647	Invitrogen	cat# A-31573; RRID:AB_2536183
chicken Alexa 488	Jackson ImmunoResearch	cat# 703-545-1551
sheep Alexa 594	Invitrogen	cat# A-11016; RRID:AB_2534083
rat Alexa 488	Invitrogen	cat# A-21208; RRID:AB_2535794
rabbit HRP	GE Healthcare	cat# NA9340V; RRID:AB_772191
mouse HRP	GE Healthcare	cat# NA931; RRID:AB_772210
Chemicals, Peptides, and Recombinant Proteins		
DL-Norepinephrine hydrochloride	Sigma-aldrich	cat# A7256
Phentolamine hydrochloride	Sigma-aldrich	cat# P7547
Atropine sulfate salt monohydrate	Sigma-aldrich	cat# A0257
Carbachol	Millipore	cat# 212385
(-)-Isoproterenol hydrochloride	Sigma-aldrich	cat# I6504
(±)-Propranolol hydrochloride	Sigma-aldrich	cat# P0884
R(-)-Denopamine	Sigma-aldrich	cat# D7815
(±)-Metoprolol (+)-tartrate salt	Sigma-aldrich	cat# M5391
Salbutamol	Sigma-aldrich	cat# S8260
ICI-118,551 hydrochloride	Sigma-aldrich	cat# I127
Experimental Models: Cell Lines		
MeWo	Sigma-Aldrich	cat# 93082609
human bronchial smooth muscle cells	ScienCell	cat#3400
B16F10	ATCC	CRL-6475, RRID:CVCL_0159
human dermal lymphatic endothelial cells	Group Prof. Detmar	N/A

(Continued on next page)

Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER
Experimental Models: Organisms/Strains		
Prox-1 GFP	Dr. Young-Kwon Hong, University of Southern California	I. Choi et al., 2011
ChAT GFP	Jackson Laboratories	Strain No 007902
Cx40 ^{BAC} -GCaMP2	Dr. Michael Kotlikoff, Cornell University and the CHROMus mouse resource	R24HL120847
C57BL/6J	Jackson Laboratories	Strain No 000664
Oligonucleotides		
	See Table S4 for used primers	N/A
Software and Algorithms		
Quantification algorithms	For full MATLAB codes please see STAR Methods .	N/A

CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Michael Detmar (michael.detmar@pharma.ethz.ch).

EXPERIMENTAL MODELS AND SUBJECT DETAILS

Cell culture

Primary male human LECs ([Hirakawa et al., 2003](#)) were cultured under standard culture conditions (37°C and 5% CO₂) on collagen (Advanced BioMatrix) coated dishes (50 µg/mL) in EBM medium (Lonza) containing 20% FBS (GIBCO), 1% penicillin/streptomycin (GIBCO), 2 mmol/L L-glutamine (GIBCO), 25 µmol/mL cAMP (Sigma-Aldrich) and 10 µg/mL hydrocortisone (Sigma-Aldrich).

Human bronchial smooth muscle cells were purchased from ScienCell and cultured under standard conditions in Smooth Muscle Cell Medium 2 containing the Supplemental Mix (both PromoCell). MeWo cells (Sigma Aldrich) were cultured under standard culture conditions in DMEM containing pyruvate (GIBCO) and 10% FBS.

Mice

Mice were group housed in IVC cages containing a mouse house and environmental enrichment under pathogen-free conditions. Mice were not involved in previous procedures. Mice of both genders were used at the age of 7-20 weeks for organ collection and 7-12 weeks for *in vivo* imaging. *Prox1-GFP* mice on the C57BL/6J background, used for clear visualization of LVs in whole mounts and tumor studies, were a kind gift of Dr. Young-Kwon Hong, University of Southern California ([Choi et al., 2011](#)). *B6.Cg-Tg(RP23-268L19-EGFP)2Mik/J* (ChAT-GFP, strain number 007902) mice and *C57BL/6J* mice were purchased from Jackson Laboratories, Bar Harbor, ME. *Cx40^{BAC}-GCaMP2* mice ([Tallini et al., 2007](#)) were obtained from Dr. Michael Kotlikoff, Cornell University and the CHROMus mouse resource (R24HL120847).

For organ collection at the end of imaging, mice were euthanized with an overdose of anesthesia (160 mg/kg ketamine; 0.4 mg/kg medetomidine) followed by cervical dislocation and opening of the chest cavity. All experiments were approved by Kantonales Veterinäramt Zurich (protocols: 11/2012, 12/2015).

METHOD DETAILS

LEC treatment *in vitro*

LECs were incubated in EBM containing 1% FBS overnight (starvation medium). Compounds of interest ([Table S1](#)) were added in starvation medium. Antagonists were added 30 min before treatment of the cells with the corresponding agonist. 15 min after agonist addition, total protein was extracted and protein concentration was measured using a Pierce BCA assay (ThermoFisher).

Western blotting

Equal amounts of denatured protein were loaded per slot, using pre-cast NuPage gels (ThermoFisher). A Mini Gel Tank system was used to run and blot the gels (ThermoFisher) according to manufacturer's instructions. After blocking the membranes, protein was detected using specific primary antibodies ([Table S2](#)) and HRP-conjugated secondary antibodies ([Table S3](#)). ECL was used for visualization using an Agfa Curix Developer. Quantification of phosphorylation was performed using FIJI ([Schindelin et al., 2012](#)).

RNA isolation and qPCR

For RNA extraction, the NucleoSpin RNA kit from Macherey-Nagel was used according to the manufacturer's instructions. RNA was reverse transcribed with the High Capacity cDNA kit (Applied Biosystems/ThermoFisher). qPCR analyses were done in triplicate using SYBRGreen (Roche) on a 7900 HT Fast Real-Time PCR System (Applied Biosystems). The following genes were analyzed: *ADRA1A*, *ADRA1B*, *ADRA1d*, *ADRA2A*, *ADRA2B*, *ADRA2C*, *ADRB1*, *ADRB2*, *ADRB3*, *ChRM1*, *ChRM2*, *ChRM3*, *ChRM4* and *ChRM5* (for primers see [Table S4](#)). For normalization, *GAPDH* expression was analyzed.

Tissue digestion and FACS analysis

For FACS analysis of LECs, the flank collectors were dissected, minced, and digested in a collagenase solution [530 u/mL Collagenase II (Worthington), 0.744 u/mL Elastase (Worthington), 5 u/mL DNase I (Roche)] for 70 min at 37°C. The digested tissue was passed through a cell strainer, washed and labeled with antibodies (see [Tables S2](#) and [S3](#)) for 30 min at 4°C, followed by incubation of secondary antibody for 30 min at 4°C. Cells were stained for Zombie-NIR used for life/dead discrimination. After washing, cells were resuspended in FACS buffer for acquisition. Data were acquired on a FACS ARIA (BD) and analyzed using FlowJo v10.2 software (Treestar Inc.).

Imaging of flank collector contractions

The contractility of the flank collecting LV was analyzed as described ([Chong et al., 2016](#)). In brief, mice were anesthetized using a combination of xylazine (20 mg/kg, Rompun® 2%, Provet), ketamine (100 mg/kg, Ketazol®, Graeub) and acepromazine (3 mg/kg, Prequillan®, Arovet) or with isoflurane (2.5%, Attane®, Provet). After loss of a detectable response to toe pinches, mice were fixed on a silicone dish and the flank collector was exposed. Tissue was kept moist by applying warm PBS and mice were kept on a heating pad during the imaging. Cannulation of the inguinal lymph node enabled a 0.5 μL infusion of 10 μmol/L P20D680 dye into the node and thereby visualization of the flank collector. For infusion, a pump (PHD2000, Harvard Apparatus, Cambridge, MA) was used, connected to a custom-made catheter using polyethylene PE-10 tubing (SCI, Lake Havasu City, AZ) and a 30-g needle. After stabilization of the contractility, videos of 8 min (9 min for the specificity testing) duration were acquired. Local topical treatment of the flank collector was done at time point 2 min (compound information see [Table S1](#)). For the specificity testing, the second treatment was topically added at time point 4 min. Videos were acquired using either a Zeiss StereoLumar.V12 stereomicroscope with AxioVision (Carl Zeiss, Feldbach, Switzerland) software and a Photometrics Evolve 512 camera or a Zeiss AxioZoom.V16 fluorescence zoom microscope with Zen Pro (Carl Zeiss) software and a QImaging OptiMOS sCMOS camera (QImaging, Surrey, Canada) in combination with a light-emitting diode illumination system pE-4000 (CoolLED Ltd, Andover, UK) and filters for Cy5 (Carl Zeiss). The contractility was quantified using a previously described algorithm ([Chong et al., 2016](#)).

Mice were allocated randomly to a specific treatment. Group size was evaluated after first pilot studies using power-analysis. Exclusion criteria for an infused vessel were: complete coverage with fat or visible luminal obstruction of the vessel induced by the infusion by e.g., blood coagulates.

In vivo Ca²⁺ imaging

Flank collectors of *Cx40^{BAC}-GCaMP2* mice were exposed. Infusion of either 0.5 μL of 10 μmol/L P20D680 tracer or a mixture of tracer with agonists was performed into the inguinal lymph node. During infusion, a video of the flank collector was acquired using a custom built 2-photon laser-scanning microscope. For quantification of a possible Ca²⁺ peak, mean intensity of the GCaMP2 signal was measured over the vessel wall during the video ([Figure 5A](#)). Specific compounds were randomly allocated to mice.

In vitro Ca²⁺ imaging

The Fluo8H, AM (AAT Bioquest, cat-no. 21091) was used according to the manufacturer's instructions. In brief, Fluo8H stock solution (1.14 mol/L) was prepared by adding 1.5% Pluronic and 2.4% DMSO in PBS. Cells were incubated in EBM containing 1% FBS for 4h, Fluo8H 1:500 in starvation medium was added and cells were incubated for 30 min at 37°C, followed by 10 min at room temperature. Cells were washed twice with Live Cell Imaging Solution (Molecular Probes). Imaging of cells was performed in Live Cell Imaging Solution; compounds were added in the same volume of live cell imaging solution as the cells were cultured in, using 2x treatment solutions. Imaging was performed using a fluorescent microscope (Nikon Eclipse Ti-E, Tokyo, Japan) in combination with a Hamamatsu ORCA-Flash4.0 CCD camera (Hamamatsu, Japan). FITC fluorescence videos were acquired and quantified as stated below.

Whole mount stainings

Vessels or pieces of small intestine were dissected out of 7- to 20-week-old mice of either gender. The small intestine samples were cut open longitudinally to give access to villi. Samples were fixed with 4% PFA for 2h at 4°C or for 1h in 100% methanol at 4°C. After washing in PBS, tissues were incubated in 0.2% BSA, 5% donkey serum, 0.1% Triton-X in PBS for 3h at 4°C. Primary antibodies were incubated over night at 4°C. After washing in 0.1% Triton X in PBS, secondary antibodies were applied in 0.1% Triton X in PBS for 2h at room temperature or at 4°C for the small intestine. After several washes in PBS, tissues were mounted using Vectashield (Vector) for confocal imaging. The antibodies used are listed in [Tables S2](#) and [S3](#). Whole mount pictures were acquired using a LSM 780 FCS confocal microscope and ZEN software (Zeiss, Jena, Germany) and processed with FIJI. Positive control stainings

for the neurotransmitter receptors are as follows: α_1 -positive enterocytes are visible in [Figure 1C](#), β_1 -positive heart tissue is provided in [Figure S1D](#), β_2 -positive blood vessels are visible in [Figure 1D](#) and M_2 -positive liver tissue is provided in [Figure S1E](#).

B16F10 melanoma studies

Male *C57BL/6J* mice of 7 weeks were bilaterally injected intradermally with 10^5 B16F10 tumor cells in the flank skin. Tumor growth was monitored by size measurements using a caliper. When tumors reached 12 mm in one dimension, tumor-draining LVs were imaged *in vivo*. Mice were randomly assigned to compound treatments.

QUANTIFICATION AND STATISTICAL ANALYSIS

Quantification of innervation

Overall length of nerves covering vessels was measured manually in FIJI and divided by the length of the LV.

Statistical analyses

All statistical analyses were performed using the GraphPad Prism 5 software. Data are shown as means \pm SD, information on exact value and representation of *n* can be found in the respective figure legends. For comparison of more than two groups, one-way ANOVA with Bonferroni post-test was used for normally distributed data. For quantifications from *in vivo* flank imaging, a paired two-tailed Student's *t* test was used for normal distributed data. A Mann-Whitney test was used for the comparison of two groups if the data was non-normal distributed, tested by a Shapiro-Wilk test. A *p* value of *p* < 0.05 was considered significant (*p* < 0.05 *, *p* \leq 0.01 **, *p* \leq 0.001***).

DATA AND SOFTWARE AVAILABILITY

Quantification of low frequency contractions

Experiments performed with isoflurane resulted in a much lower contraction activity which could not be analyzed with the previously developed algorithm ([Chong et al., 2016](#)). Therefore, a new algorithm was developed ([Figure S4](#)). In order to determine the characteristics of the peaks (i.e., amplitudes and periodicity) of the measured fluorescence signal, a MATLAB (MathWorks Inc., Natick MA, USA) function with a graphical user interface was developed that performs the signal processing in a semi-automatically manner. The user selects the peaks (α_i) of the fluorescence signal manually and based on these selected values, the absolute amplitudes (A_j^{abs}), relative amplitudes (A_j^{rel}) and the frequency (F_j) are then calculated automatically. These parameters were calculated according to the formulas

$$A_j^{abs} = |M_j - \alpha_{i+1}|, \quad (1)$$

$$A_j^{rel} = 100 \times \left(A_j^{abs} / M_j \right), \quad (2)$$

$$F_j = 1/P_j, \quad (3)$$

with M_j the baseline value of each peak according to $M_j = \alpha_i + (\alpha_{i+2} - \alpha_i)/2$ in case of $\alpha_{i+2} > \alpha_i$ or $M_j = \alpha_{i+2} + (\alpha_i - \alpha_{i+2})/2$ in case of $\alpha_{i+2} < \alpha_i$, and P_j the time-difference between the two peaks α_i and α_{i+1} with the first peak assigned to $i = 1$ and the second to $i = 2$ (and so on). For a visualization of the mathematical nomenclature used, see [Figure S4](#). The formulation ensures that the amplitude and periodicity values are also calculated correctly in case the signal contains negative peaks, or a mixture of positive and negative peaks.

MATLAB Code

```
function [Amplitude_abs, Amplitude_rel, Period, Frequency] = AnalyzePeaks_Ampl_Freq_v2(x, f_s)
% Function to analyze a time-series regarding the characteristics of the
% peaks (i.e., amplitudes, period)
%
% _____
% For Steven Proulx and Samia Bachmann, ETH.
% May 2017 | Felix Scholkmann | Version 2
%
% _____
% INPUT
% x: signal to analyze
% f_s: sampling frequency in Hz
% OUTPUT
```



```

% Amplitude_abs: absolute amplitude values (first value: median, second value: inter quantile range,
IQR)
% Amplitude_rel: relative amplitude values (first value: median, second value: inter quantile range,
IQR)
% Period: period values [in seconds] (first value: median, second value: inter quantile range, IQR)
% Frequency: frequency values [in 1/min] (first value: median, second value: inter quantile range, IQR)
% important:
% The input signal (x) must contain at least 3 peak (i.e., at least 9
% points needs to be selected from the signal). This is necessary to be
% able to calculate the period values.

```

```
t = [1 : length(x)]./f_s;
```

```

%% _____
close all
scrsz = get(0, 'ScreenSize'); figure('Position', [100 50 1300 850]);
set(gcf., 'color', 'w'); set(gcf., 'defaultaxesfontsize', 12)
subplot(2, 20, [1, 3])
% Please insert here the path to the file ``Points.png``
currentFolder = pwd;
Path = strcat(currentFolder, '\Points.png')
img = imread(Path);
image(img);
set(gca, 'XTickLabelMode', 'Manual')
set(gca, 'XTick', [])
set(gca, 'YTickLabelMode', 'Manual')
set(gca, 'YTick', [])
title('Definition', 'fontsize', 14)
fig = subplot(2, 20, [6, 20]);
plot(t, x, 'linewidth', 2, 'Color', [0.3, 0.5, 1])
axis tight
ylabel('Intensity'); xlabel('Time [s]')
title('- > Please select the points (\alpha_{i}, \alpha_{i+1}, \alpha_{i+2}, i = 1, 2, ..., \it
{N})', 'fontsize', 14)
% Get the points from the figure
disp('\to Please select the points in the figure. The press ``Enter``')
[x, y] = getpts(fig);
hold on
vline_red(x(2:3:end))
vline_black(x(1:3:end))
vline_black(x(3:3:end))
n = length(x);
% Calculate the amplitudes
j = 1;
for i = 1:3:n-2;
    if y(i+2) > y(i)
        M(j) = (y(i+2)-y(i))/2 + y(i);
        A_abs(j) = abs(M(j) - y(i+1));
        A_rel(j) = 100 * (A_abs(j)/M(j));
        j = j+1;
    else
        M(j) = (y(i)-y(i+2))/2 + y(i+2);
        A_abs(j) = abs(M(j) - y(i+1));
        A_rel(j) = 100 * (A_abs(j)/M(j));
        j = j+1;
    end
end

```

```

        end
    end
    plot(x(2:3:end),M,'ob')
    plot(x(1:3:end),y(1:3:end),'ok')
    plot(x(3:3:end),y(3:3:end),'ok')
    plot(x(2:3:end),y(2:3:end),'or')
    % Calculate the periods
    j = 1;
    for i = 1:3:n-4;
        P(j) = x(i+4)-x(i+1);
        j = j+1;
    end
    F = (1./(P))*60; % Frequency [1/min]
    subplot(2,20,[21:23])
    bar(A_abs,'b');
    ylabel('Amplitude (A_{abs}) [absolute]'); xlabel('Peak number')
    title({'Amplitude values', '[absolute]'}, 'fontsize', 14)
    xlim([0,length(A_abs)+1])
    subplot(2,20,[25:27])
    bar(A_rel,'b');
    ylabel('Amplitude (A_{rel}) [%]'); xlabel('Peak number')
    title({'Amplitude values', '[%]'}, 'fontsize', 14)
    xlim([0,length(A_rel)+1])
    subplot(2,20,[29:31])
    bar(F,'r');
    ylabel('Frequency [1/min]'); xlabel('Peak number')
    title({'Frequency values', '[1/min]'}, 'fontsize', 14)
    xlim([0,length(F)+1])
    subplot(2,20,35:36)
    errorbar(median(A_rel),iqr(A_rel)/2,'k');
    hold on
    plot(median(A_rel),'k.','markersize',30)
    ylabel('Amplitude [%] (median +- IQR)')
    set(gca, 'XTickLabelMode', 'Manual')
    set(gca, 'XTick', [])
    title(['A_{rel} = ' num2str(median(A_rel)), ' (' num2str(iqr(A_rel)), ') %']);
    subplot(2,20,39:40)
    errorbar(median(F),iqr(F)/2,'k');
    hold on
    plot(median(F),'k.','markersize',30)
    ylabel('Frequency [1/min] (median +- IQR)')
    set(gca, 'XTickLabelMode', 'Manual')
    set(gca, 'XTick', [])
    title(['F = ' num2str(median(F)), ' (' num2str(iqr(F)), ') 1/min']);
    % Generate output data
    Amplitude_abs = [median(A_abs), iqr(A_abs)];
    Amplitude_rel = [median(A_rel), iqr(A_rel)];
    Period = [median(P), iqr(P)];
    Frequency = [median(F), iqr(F)];

```

Quantification of *in vitro* Ca²⁺ imaging

Pre-processing. An important step to restore images is denoising, which consists of reducing noise present in images but at the same time preserving objects' features such as edges, texture, etc. In this context, we have opted to use the Block-Matching and 3D filtering (BM3D) method which has shown its denoising capabilities in different datasets by preserving finer image structures (Dabov et al., 2007; Danielyan et al., 2014).

Cell Detection. Once the image has been pre-processed, the next step is to detect the foreground objects, i.e., cells. To do this, we use the Otsu thresholding method (Otsu, 1979), however instead of using a single threshold value, we adopt a two-level threshold to detect the nucleus and the cytoplasm as joint single objects.

After thresholding, small objects are further removed since they usually correspond to non-relevant objects, such as air-bubbles. In addition, since some cells may spatially be close to each other, they can end up being merged after thresholding. To cope with this issue, we use the Watershed Transform. More precisely, each segmented object is represented as a basin and the detected Watershed Lines with the distance transform are used to separate the objects.

Cell Linking. Cells are detected at each single frame separately. In order to analyze how a cell activates over time, the detected cells are first linked together over time using the Kalman filter (Kalman, 1960).

The Kalman filter comprises two steps: prediction and update. In the prediction step, the Kalman filter predicts the current state of the system, based on the past, through a dynamic model. In the update step, the status of the system is corrected based on the observed variables. In our case, the assignment/linking of a given cell to a specific track is based on the area of overlap of the segmented cells between consecutive frames. The Kalman Filter therefore seeks to maximize the area of overlap between cell detections over time. To obtain more robust predictions at time t , our system averages the bounding-box of the detected cells over the last four frames. To remove false positives such as air-bubbles, our system discards objects with large location changes over time. Also, short tracks are discarded for further processing. In our case, we impose the constraint that tracks shall cover at least 75% of the total length of the input video.

Event Detection. Our event detection module is versatile and besides detecting which cells got activated, it can also find out when exactly a given activation starts and ends over time. The module starts first by smoothing the cell's intensity signal (the cell-intensity track) to reduce noise. Smoothing is done with a moving average filter. Note that the smoothing step is optional and the raw signal can be directly used if so desired.

Next, the cell signals are normalized to compensate for effects such as photobleaching and also to facilitate the detection of cell activations in the subsequent steps. The normalization of a cells' signal is done by computing the average intensity μ_i , of the detected cell at the frames before the injection time t_{inj} (see Equation 1).

$$\mu_i = \frac{1}{t_{inj}} \sum_{t=0}^{t=t_{inj}} f(x_i^t) \quad (\text{Equation 1})$$

where $f(\cdot)$ is the mean intensity value of the collection of pixels x_i^t belonging to the detected cell i at time t .

The computed mean value μ_i of cell i is used to normalize the signal by dividing it into the cell signal values at all later times (see Equation 2).

$$\hat{f}(x_i) = \frac{1}{\mu_i} f(x_i^t); t = \{t = t_{inj} + 1, \dots, N\} \quad (\text{Equation 2})$$

where \hat{f} is the normalized cells' signal and N corresponds to the total number of frames. Figure S3B shows an example of cell signals before (upper graph) and after normalization (lower graph). Detected activations above the threshold value (displayed as a black dashed line) are shaded, see next section of details of the detection algorithm.

To detect cell-activations, the normalized cell signals are thresholded. More precisely, signal values larger than a threshold value are considered as activations and values lower than the threshold are set to zero. We set this threshold value empirically to be 1.25. Note that this value can be tuned, if more detections are targeted, one can increase this value. It is important though to set this value in a conservative manner. An example of cell activation is displayed in yellow in the left graph of Figure S3C. The thresholding step is applied for all cell signals, in such a way that all activations can be mapped as an activation map. In Figure S3C, each row represents a single cell signal and its mean-intensity pixel values are displayed for each frame along the x axis. After thresholding the cell signals on the normalized cell signals (middle graph in Figure S3C), a binary activation map is obtained in which cell activations are displayed as white stripes (right graph, Figure S3C). The signal values where the cell did not activate are set to zero. In addition, to remove potential false negatives, our method first fills short gaps inside an event and also prunes short cell activations to remove false positives. In our experiments, both values were empirically set to five frames.

One important property of our method is that it detects where a cell activation starts and ends. This is done by computing the first order derivative over the binary activated signal and by looking respectively for the indexes where the derivative is respectively positive or negative (i.e., +1 or -1).

MATLAB Code (2 parts)

```
function kalmanMultiObjectTracking(videoName)
% KalmanMultiObjectTracking: this function detects a set of cells and
% tracks them over time. Cell detection involves image pre-processing where
% the original video frames are enhanced, e.g., denoising. The cell
% detections are mapped into cell-tracks, i.e., a set of temporal
% bounding-boxes for each detected cell.
%
```

```

% Input:
% videoName, e.g., videoName = 'hu_NE1uM_2.avi';
%
% Output:
% tracksHist.mat MATLAB file containing the cell-tracks.
opts = getDefaultParameters(); %# reading configuration parameters.
pathInputVideo = fullfile(opts.videoPath, videoName);
digitStr = '%d-';
strAllMotionDigits = repmat(digitStr, 1, numel(opts.motionNoise));
motionNoiseStr = sprintf(strAllMotionDigits(1:end-1), opts.motionNoise);
strAllInitErrorDigits = repmat(digitStr, 1, numel(opts.initialEstimateError));
initErrorStr = sprintf(strAllInitErrorDigits(1:end-1), opts.initialEstimateError);
paramsDir = sprintf('%s_LRate%2.2f_visi%2.2f_age%d_inviv%d_initErr%s_motNoise%s_measNoise%d_
%s_cSize%d', ...
    opts.motionModel, opts.learningRate, opts.trackVisibility, opts.ageThreshold, ...
    opts.invisibleForTooLong, initErrorStr, motionNoiseStr, opts.measurementNoise, opts.-
    thresh.type, opts.cell.size);
[videosPath, inVideoname, ~] = fileparts(pathInputVideo);
videoNameOutputTrackRGB = sprintf('%s_track_rgb.avi', inVideoname);
videoNameOutputTrackProc = sprintf('%s_track_proc.avi', inVideoname);
videoNameOutputTrackBW = sprintf('%s_track_bw.avi', inVideoname);
videoNameOutputBW = sprintf('%s_bw.avi', inVideoname);
tracksDir = fullfile(videosPath, 'tracks_kalman_sel');
tracksDirVideo = fullfile(tracksDir, inVideoname, paramsDir);
opts.framesOrigDir = fullfile(tracksDir, inVideoname, 'frames');
opts.framesProcDir = fullfile(tracksDirVideo, 'frameProc');
opts.maskDir = fullfile(tracksDirVideo, 'frameMasks');
opts.maskDirTracks = fullfile(tracksDirVideo, 'frameMasksTracks');
opts.framesOrigTracksDir = fullfile(tracksDirVideo, 'framesOrigTracks');
opts.framesProcTracksDir = fullfile(tracksDirVideo, 'framesProcTracks');
opts.maskDirTracksAll = fullfile(tracksDirVideo, 'frameMasksTracksAll');
opts.framesOrigTracksDirAll = fullfile(tracksDirVideo, 'framesOrigTracksAll');
createVideoDirs(tracksDirVideo, opts); %# creating output directories
videoObjTrackRGB = VideoWriter(fullfile(tracksDirVideo, videoNameOutputTrackRGB),
    'Uncompressed AVI');
videoObjTrackProc = VideoWriter(fullfile(tracksDirVideo, videoNameOutputTrackProc),
    'Uncompressed AVI');
videoObjTrackBW = VideoWriter(fullfile(tracksDirVideo, videoNameOutputTrackBW), 'Uncompressed
    AVI');
videoObjBW = VideoWriter(fullfile(tracksDirVideo, videoNameOutputBW), 'Uncompressed AVI');
fprintf('video: %s\n', (fullfile(tracksDirVideo, videoNameOutputTrackRGB)));
tracksObj = fullfile(tracksDirVideo, sprintf('%s.mat', inVideoname));
obj = setupSystemObjects(pathInputVideo, opts);
tracks = initializeTracks(); %# Create an empty array of tracks.
tracksHist = initializeTracksHistory();
nextId = 1; % ID of the next track
noFrames = 0;
if ~exist(tracksObj, 'file')
    %# Detect moving objects, and track them across video frames.
    while ~isDone(obj.reader)
        noFrames = noFrames + 1;
        fprintf('# Processing frame: %d\n', noFrames);
        frame = readFrame(obj);
        %# image pre-processing and per-frame cell detection.
        [centroids, bboxes, mask, frameDen] = detectObjects(frame, opts);
        %# predict cell location.
        [tracks, tracksHist] = predictNewLocationsOfTracks(tracks, tracksHist, noFrames);
        %# link cells over time.
    end
end

```

```

    [assignments, unassignedTracks, unassignedDetections] = ...
detectionToTrackAssignment(tracks, centroids, bboxes, opts);
    %# update existing cell-tracks.
    [tracks, tracksHist] = updateAssignedTracks(tracks, tracksHist, assignments, centroids,
bboxes, noFrames);
    [tracks, tracksHist] = updateUnassignedTracks(tracks, tracksHist, unassignedTracks);
    %# delete short tracks.
    [tracks, tracksHist] = deleteLostTracks(tracks, tracksHist, opts);
    %# create new tracks.
    [tracks, tracksHist, nextId] = createNewTracks(tracks, tracksHist, centroids, bboxes,
unassignedDetections, nextId, noFrames);
    %# saving tracks to disk.
    dumpTrackingResults(frame, mask, tracks, tracksHist, tracksDirVideo, videoObjTrackRGB,
videoObjTrackBW, videoObjBW, noFrames, opts);
end
opts.totNoFrames = noFrames;
tracksHist = updateEmptyTracks(tracksHist, opts); %# check empty tracks frames and reduce its
totalVisibleCount
save(tracksObj, 'tracksHist', '-v7.3'); %# save to disk the detected tracks.
else
    fprintf('# Loading tracks\n');
    tracksHist = load(tracksObj);
    tracksHist = tracksHist.tracksHist;
    reset(obj.reader);
    while ~isDone(obj.reader)
        frame = readFrame(obj);
        noFrames = noFrames + 1;
    end
    fprintf('# No of. frames: %d\n', noFrames);
end
opts.totNoFrames = noFrames;
if ~isfield(tracksHist, 'numEmptyMask')
    tracksHist = updateEmptyTracks(tracksHist, opts);
end
open(videoObjTrackRGB); %# create RGB video with cell tracks.
open(videoObjTrackProc); %# create RGB video for cell tracks (segmented/cells + bounding boxes).
open(videoObjTrackBW); %# create BW video for cell tracks (segmented/detected cells + bounding
boxes).
open(videoObjBW); %# create BW video file for cell detections (segmented/detected cells as binarys
masks).
noFrames = 0;
for ff = 1:opts.totNoFrames
    noFrames = noFrames + 1;
    frameOrigFile = fullfile(opts.framesOrigDir, sprintf('frame_%05d.tif', ff));
    frameDenFile = fullfile(opts.framesProcDir, sprintf('frame_%05d.tif', ff));
    frameMaskFile = fullfile(opts.maskDir, sprintf('frame_%05d.tif', ff));
    fprintf('#Processing frame: %d\n', noFrames);
    if exist(frameOrigFile, 'file') && exist(frameDenFile, 'file') && exist(frameMaskFile, 'file')
        frameOrig = imread(frameOrigFile);
        frameDen = imread(frameDenFile);
        frameMask = imread(frameMaskFile);
        %# storing into disk each frame overlaid with the cell-tracks.
        saveTrackingResults(frameOrig, frameDen, frameMask, tracksHist, videoObjTrackRGB,
videoObjTrackProc, videoObjTrackBW, videoObjBW, ff, opts);
    end
end
close(videoObjTrackRGB); %# closing videoFile.
close(videoObjTrackProc); %# closing videoFile.

```

```

close(videoObjTrackBW); %# closing videoFile.
close(videoObjBW); %# closing videoFile.
end
function tracksThresh2Activations(videoName)
% tracksThresh2Activations: given the extracted cell-tracks of a
% given video (linked cells over time), this function detects which of
% the cells got activated. For that purposes, the mean pixel intensity
% of a segmented cell is computed at each frame. If the mean pixel
% intensity value is higher than a threshold, then an activation is fired.
% The original intensity values can also be smoothed to remove noise and if
% the interval of the activation is too short, then a false-positive activation
% is detected.
%
% Input:
% videoName, e.g., videoName = 'hu_NE1uM_2.avi';
%
% Output:
% a set of figures are stored into disk, containing a
% matrix-representation of the activated cell-signals.
addpath(genpath('./3rd/overlay/'));
%# reading default parameters, e.g., location of detected cells frames.
opts = getDefaultParameters();
if ~isfield(opts, 'endThreatment')
    opts.endThreatment = 10;
end
if ~isfield(opts, 'signal')
    opts.signal = [];
end
if ~isfield(opts.signal, 'Type')
    opts.signal.Type = 'orig';
end
%# smoothing type for the cell signal.
if ~isfield(opts.signal, 'smoothType')
    opts.signal.smoothType = 'box'; % 'golay', 'gauss', 'box'
end
%# threshold value
if ~isfield(opts.signal, 'threshVal')
    opts.signal.threshVal = 1.25;
end
%# minimum duration of cell-activation
if ~isfield(opts.signal, 'pruneEventDuration')
    opts.signal.pruneEventDuration = 5;
end
if ~isfield(opts.signal, 'fillGaps')
    opts.signal.fillGaps = true;
end
if ~isfield(opts.signal, 'fillGapsSze')
    opts.signal.fillGapsSze = 5;
end
[videosPath, videoNameInput, ~] = fileparts(videoName);
digitStr = '%d-';
strAllMotionDigits = repmat(digitStr, 1, numel(opts.motionNoise));
motionNoiseStr = sprintf(strAllMotionDigits(1:end-1), opts.motionNoise);
strAllInitErrorDigits = repmat(digitStr, 1, numel(opts.initialEstimateError));
initErrorStr = sprintf(strAllInitErrorDigits(1:end-1), opts.initialEstimateError);
paramsDir = sprintf('%s_LRate%2.2f_visi%2.2f_age%d_in vib%d_initErr%s_motNoise%s_measNoise%d_
%s_cSze%d', ...
    opts.motionModel, opts.learningRate, opts.trackVisibility, opts.ageThreshold, ...

```

```

    opts.invisibleForTooLong, initErrorStr, motionNoiseStr, opts.measurementNoise,
    opts.thresh.type, opts.cell.size);
%# creating subdirectories where results are going to be stored.
tracksDir = fullfile(videosPath, 'tracks_paper');
tracksDirVideo = fullfile(tracksDir, videoNameInput, paramsDir);
signalDir = fullfile(tracksDirVideo, 'signal');
signalThreshDir = fullfile(signalDir, 'thresh');
signalEventsDir = fullfile(signalDir, 'events');
if ~exist(signalEventsDir, 'dir'), mkdir(signalEventsDir); end
tracksFile = fullfile(tracksDirVideo, 'validTracksIdx.mat');
validTracksIdx = load(tracksFile);
validTracksIdx = validTracksIdx.validTracksIdx;
%# loading detected cell-tracks
if strcmpi(opts.signal.Type, 'orig')
    threshSubDir = fullfile(signalThreshDir, 'origdata');
    signalMeanFile = fullfile(tracksDirVideo, 'signalMean.mat');
    signalMean = load(signalMeanFile);
    signalMean = signalMean.signalMean;
    signalMean(~validTracksIdx, :) = [];
    signalArray = signalMean;
    if ~exist(threshSubDir, 'dir'), error('Unknown directory: %s\n', threshSubDir); end
    signalEventsDir = fullfile(signalEventsDir, 'origdata');
    if ~exist(signalEventsDir, 'dir'), mkdir(signalEventsDir); end
    signalTypeStr = opts.signal.Type;
    tracksIdx = find(validTracksIdx);
elseif strcmpi(opts.signal.Type, 'smooth')
    fprintf('# Loading %s data\n', opts.signal.smoothType);
    signalSmoothFile = fullfile(signalDir, 'smooth', sprintf('tracksSummary_%s.mat',
    opts.signal.smoothType));
    signalMeanSmooth = load(signalSmoothFile);
    signalMeanSmooth = signalMeanSmooth.signalMeanSmooth;
    signalMeanSmooth(~validTracksIdx, :) = [];
    signalArray = signalMeanSmooth;
    threshSubDir = fullfile(signalThreshDir, opts.signal.smoothType);
    if ~exist(threshSubDir, 'dir'), error('Unknown directory: %s\n', threshSubDir); end
    signalEventsDir = fullfile(signalEventsDir, opts.signal.smoothType);
    if ~exist(signalEventsDir, 'dir'), mkdir(signalEventsDir); end
    signalTypeStr = sprintf('%s-%s', opts.signal.Type, opts.signal.smoothType);
    tracksIdx = 1:size(signalArray, 1);
else
    error('Unkown type: %s', opts.signal.Type);
end
signalArrayNormFilename = fullfile(threshSubDir, 'signalNorm.mat');
if ~exist(signalArrayNormFilename, 'file'), error('Unknown file: %s\n', signalArrayNormFilename);
end
signalNorm = load(signalArrayNormFilename);
signalNorm = signalNorm.signalNorm;
%# thresholding for activation.
signalBinaryActivation = signalNorm > opts.signal.threshVal;
[noTracks, ~] = size(signalBinaryActivation);
%# fill possible gaps in the cell-activations
if opts.signal.fillGaps > 0
    filledSignal = signalBinaryActivation;
    for ii = 1:noTracks
        filledSignal(ii, :) = ~bwareaopen(~filledSignal(ii, :), opts.signal.fillGapsSize);
    end
    signalBinaryActivation = filledSignal;
    strFillGap = sprintf('fillSize%d', opts.signal.fillGapsSize);

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else
    strFillGap = sprintf('nofill');
end
signalActivation = zeros(size(signalBinaryActivation));
if opts.signal.pruneEventDuration > 0
    strPruneEvent = sprintf('pruned%d', opts.signal.pruneEventDuration);
else
    strPruneEvent = sprintf('nopruned');
end
end
%# analyzing cell-track duration.
stats = repmat(struct('startIdx', [], 'endIdx', [], 'duration', [], 'eventIdx', [], 'activation',
0), noTracks, 1);
for tt = 1:noTracks
    trackActivation = signalBinaryActivation(tt,:);
    trackActivation = diff([0 trackActivation 0]);
    eventStartIdx = find(trackActivation > 0);
    eventEndIdx = find(trackActivation < 0)-1;
    eventDuration = eventEndIdx - eventStartIdx + 1;
    if opts.signal.pruneEventDuration
        stringIdx = (eventDuration >= opts.signal.pruneEventDuration);
        eventStartIdx = eventStartIdx(stringIdx);
        eventEndIdx = eventEndIdx(stringIdx);
    end
    eventIdx = zeros(1,max(eventEndIdx)+1);
    eventIdx(eventStartIdx) = 1;
    eventIdx(eventEndIdx+1) = eventIdx(eventEndIdx+1)-1;
    eventIdx = find(cumsum(eventIdx));
    signalActivation(tt, eventIdx) = 2;
    signalActivation(tt, eventStartIdx) = +1;
    signalActivation(tt, eventEndIdx) = -1;
    stats(tt).startIdx = eventStartIdx;
    stats(tt).endIdx = eventEndIdx;
    stats(tt).duration = eventDuration;
    stats(tt).eventIdx = eventIdx;
    stats(tt).activation = isempty(eventIdx) == 0;
end
videoName4Plot = strrep(videoNameInput, '_', '\_');
signalImFilename = fullfile(signalEventsDir, sprintf('%s_signal_%s_label.tif', videoNameInput,
signalTypeStr));
normSignalFilename = fullfile(signalEventsDir, sprintf('%s_signal_norm.tif', videoNameInput));
activationSignalFilename = fullfile(signalEventsDir, sprintf('%s_signal_activ_%s_%s%.tif',
videoNameInput, strPruneEvent, strFillGap, signalTypeStr));
binActivationSignalFilename = fullfile(signalEventsDir, sprintf('%s_signal_bin_activ_%s_%s_
%.tif', videoNameInput, strPruneEvent, strFillGap, signalTypeStr));
signalEventsDirBox = strrep(signalEventsDir, 'origdata', 'box');
activationBoxSignalFilename = fullfile(signalEventsDirBox, sprintf('%s_signal_activ_%s_%s_
%.mat', videoNameInput, signalTypeStr, strPruneEvent, strFillGap));
activationBoxSignalFilename = strrep(activationBoxSignalFilename, 'orig', 'smooth-box');
signalActivationBox = load(activationBoxSignalFilename);
signalActivationBox = signalActivationBox.signalActivation;
signal2disk(signalArray, videoName4Plot, signalImFilename, opts);
signal2disk(signalNorm, videoName4Plot, normSignalFilename, opts);
signal2disk(signalActivation, videoName4Plot, activationSignalFilename, opts);
signal2disk(signalActivation~= 0, videoName4Plot, binActivationSignalFilename, opts);
overlaySignalFilename = fullfile(signalEventsDir, sprintf('%s_signal_overlay_%s_%s%.tif',
videoNameInput, strPruneEvent, strFillGap, signalTypeStr));
overlayNormSignalFilename = fullfile(signalEventsDir, sprintf('%s_signal_overlaynorm_%s_%s_
%.tif', videoNameInput, strPruneEvent, strFillGap, signalTypeStr));

```



```
signal2diskOverlay(signalArray, signalActivation~ = 0, videoName4Plot, overlaySignalFilename,  
opts);  
signal2diskOverlay(signalNorm, signalActivation~ = 0, videoName4Plot,  
overlayNormSignalFilename, opts);  
overlaySignalFilename = fullfile(signalEventsDir, sprintf('%s_signal_overlay_%s_%s_%s_  
box.tif', videoNameInput, strPruneEvent, strFillGap, signalTypeStr));  
overlayNormSignalFilename = fullfile(signalEventsDir, sprintf('%s_signal_overlaynorm_%s_%s_  
%s_box.tif', videoNameInput, strPruneEvent, strFillGap, signalTypeStr));  
signal2diskOverlay(signalArray, signalActivationBox~ = 0, videoName4Plot,  
overlaySignalFilename, opts);  
signal2diskOverlay(signalNorm, signalActivationBox~ = 0, videoName4Plot,  
overlayNormSignalFilename, opts);  
signalFilenameMat = fullfile(signalEventsDir, sprintf('%s_signal_activ_%s_%s%.mat',  
videoNameInput, signalTypeStr, strPruneEvent, strFillGap));  
save(signalFilenameMat, 'signalActivation');  
signalStatsFilenameMat = fullfile(signalEventsDir, sprintf('%s_stats_%s_%s%.mat',  
videoNameInput, signalTypeStr, strPruneEvent, strFillGap));  
save(signalStatsFilenameMat, 'stats');  
end
```