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Permalink
https://escholarship.org/uc/item/5496q41c

Journal
Scientific Reports, 6(1)

ISSN
2045-2322

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Publication Date
2016-10-24

DOI
10.1038/srep35791

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Peer reviewed
Repetitive Electroacupuncture Attenuates Cold-Induced Hypertension through Enkephalin in the Rostral Ventral Lateral Medulla

Min Li, Stephanie C. Tjen-A-Looi, Zhi-Ling Guo & John C. Longhurst

Acupuncture lowers blood pressure (BP) in hypertension, but mechanisms underlying its action are unclear. To simulate clinical studies, we performed electroacupuncture (EA) in unanesthetized rats with cold-induced hypertension (CIH) induced by six weeks of cold exposure (6°C). EA (0.1 – 0.4 mA, 2 Hz) was applied at ST36-37 acupoints overlying the deep peroneal nerve for 30 min twice weekly for five weeks while sham-EA was conducted with the same procedures as EA except for no electrical stimulation. Elevated BP was reduced after six sessions of EA treatment and remained low 72 hrs after EA in 18 CIH rats, but not in sham-EA (n = 12) and untreated (n = 6) CIH ones. The mRNA level of preproenkephalin in the rostral ventrolateral medulla (rVLM) 72 hr after EA was increased (n = 9), compared to the sham-EA (n = 6), untreated CIH rats (n = 6) and normotensive control animals (n = 6). Microinjection of ICI 174,864, a δ-opioid receptor antagonist, into the rVLM of EA-treated CIH rats partially reversed EA's effect on elevated BP (n = 4). Stimulation of rVLM of CIH rats treated with sham-EA using a δ-opioid agonist, DADLE, decreased BP (n = 6). These data suggest that increased enkephalin in the rVLM induced by repetitive EA contributes to BP lowering action of EA.

Approximately one in three adults has high blood pressure (BP) worldwide. Hypertension and its consequences, such as stroke and heart attacks, are enormous public health problems. The 7th Joint National Committee on Prevention, Detection, Evaluation and Treatment of High Blood Pressure report recommends lifestyle modification (i.e., non-traditional approaches) with or without pharmacological intervention. Although the effect of acupuncture on hypertension is controversial, acupuncture may offer a non-pharmacological approach to reduce high BP.

We have shown in previous studies of patients with mild to moderate hypertension that electroacupuncture (EA) applied once weekly over an eight-week course of therapy lowers peak systolic blood pressures (SBP) by 10 – 12 mmHg as measured by 24 hr ambulatory monitoring. This response appears to persist for an additional four weeks after EA is terminated. In concert with this observed prolonged action of acupuncture, several research groups have shown significant decreases in the intensity of chronic low back and shoulder pain is maintained for one to six months after completion of acupuncture treatment provided once weekly for periods ranging between six and eight weeks, compared to placebo group. Thus, acupuncture appears to reduce chronic hypertension and pain for prolonged periods of time. However, the mechanisms underlying these long-term actions are unclear.

Epidemiological data have shown that in cold regions and during cooler seasons the prevalence of hypertension and deaths from stroke and heart attack associated with hypertension are elevated. A number of studies have demonstrated that rats exposed to mild cold (6°C) for several weeks develop hypertension (cold-induced hypertension, CIH) characteristically similar to humans with essential hypertension. The CIH experimental model is a “naturally-occurring” form of hypertension, associated with chronic cold-induced stress that is...
a common cause for essential hypertension. Cold-induced hypertension is as or more clinically relevant than elevation of BP in other hypertensive models, including spontaneous hypertensive rats (SHR), DOCA-salt and renal hypertensive rats, which commonly require surgical intervention, administration of excessive doses of drugs or hormones, or genetic manipulation. Importantly, sympathetic activity is increased in CIH. Sympathetic outflow and ultimately BP are regulated through activation of sympathoexcitatory premotor neurons in the rostral ventrolateral medulla (rVLM). Acupuncture reduces sympathoexcitatory reflex-induced elevations in BP through modulation of presympathetic rVLM neuronal activity to, in turn, decrease sympathetic activity. Thus, to simulate clinically relevant hypertension due in part to sympathoexcitation, the conscious CIH rat model was used in the present study to investigate the effect of repeated EA on BP and the mechanisms underlying EAs action in a relevant brain stem region, the rVLM.

The opioid system in the rVLM contributes to the EA-modulatory actions on reflex hypertension in anesthetized animals. Despite strong evidence showing that acupuncture influences short-term hypertensive reflexes, it is uncertain if opioids in the rVLM participate in the BP lowering actions of repetitive EA in chronic hypertension. As such, in the present study, the clinically relevant CIH model was used to examine: 1) the ability of repetitive EA to reduce sustained elevations in BP; 2) central mechanisms underlying acupuncture’s hypotensive actions (Fig. 1). We hypothesized that repetitive EA lowers CIH in conscious animals through the opioid system in the rVLM.

Results

Effect of repetitive EA treatment on cold-induced hypertension. Cold exposure increased both systolic and diastolic blood pressure (SBP, DBP) within four weeks and induced sustained hypertension by six weeks (Fig. 2, panels A and B). Blood pressures of rats in the untreated hypertensive and sham-EA groups remained elevated at the end of 11 weeks. In contrast, EA reduced both the elevated SBP and DBP after six sessions; the BPs remained low throughout the period of treatment relative to the sham treated group. Although SBP and DBP of EA-treated CIH rats post EA were higher than those observed immediately after termination of EA, they were still reduced three days post EA relative to the sham-EA control (Fig. 2, panels A and B). The BP of the normotensive rats remained stable throughout the 11 week period. Cold exposure increased heart rate (HR) within four weeks and remained unchanged in the untreated hypertensive, sham-EA and EA groups throughout the remainder of the 11 week period and for three days afterwards (Fig. 2C).

Preproenkephalin mRNA expression 72 hours after EA. The rVLM preproenkephalin mRNA level was elevated 72 hr following application of 11 sessions of EA treatment in the hypertensive animals (Fig. 3). Thus, the preproenkephalin mRNA was significantly higher 72 hour after EA (14.3 ± 1) compared to the sham-EA (4.3 ± 0.5), untreated hypertensive (4.3 ± 0.6) and normotensive rats (3.8 ± 0.9).

Opioid receptor antagonism in EA-treated CIH rats. In a preliminary study we observed that bilateral microinjection of 1 mM naloxone (50 nl; a non-specific opioid receptor antagonist) into the rVLM increased SBP, DBP and mean blood pressure (MBP) in two EA-treated hypertensive rats (140 to 146, 77 to 84, and 98 to 105 mmHg in one rat, and 132 to 144, 63 to 70, and 90 to 97 mmHg in a second rat). Heart rate was not influenced by naloxone in either rat (447 to 451, and 465 to 467 bpm, respectively). To investigate the role of the δ-opioid receptor in the EA hypotensive effect in animals with sustained hypertension, we employed specific δ-opioid receptor antagonism with ICI 174,864 (1 mM in 50 nl). We noted a slight increase in BP (−10 mmHg; Fig. 4A), but not heart rate after the initial unilateral injection. Bilateral blockade of δ-opioid receptors in the rVLM partially reversed EAs action on systolic, mean and diastolic BPs in four additional animals (Fig. 4). Heart rates were not influenced by ICI 174,864. The onset and duration of the responses following the initial unilateral injection were 10 s and 70 min, respectively, with a peak effect observed at 3 min (Fig. 4A). Microinjection of the same volume of vehicles for naloxone and ICI 174,864 in EA treated hypertensive rats did not influence BP or heart rate (n = 4).

Delta-opioid receptor stimulation in sham-EA CIH rats. The δ-opioid receptor agonist, DADLE (10 nM in 50 nl), microinjected bilaterally into the rVLM of six sham-EA CIH rats decreased SBP, but did not alter
DBP, MBP (Fig. 5) or heart rate. We noted only a small reduction in SBP (~5 mmHg) after unilateral injection. The onset (after initial unilateral injection) and duration of the responses to DADLE were 15 and 300 s, while the peak onset (after initial unilateral injection) and duration of the responses to DADLE were 15 and 300 s, while the peak
effect was observed at 140 s. Microinjection of same volume of normal saline into the rVLM did not influence BP or heart rate in four of these six rats.

**Delta-opioid receptor stimulation and blockade in normotensive rats.** Bilateral microinjection of ICI 174,864 (1 mM in 50 nl) into the rVLM of three normotensive rats did not change BP. The recordings of SBP, DBP and MBP were the same before and after the microinjection of ICI 174,864 into the rVLM of the normotensive rats, noted as 109 ± 8, 65 ± 4 and 80 ± 4 mmHg. Three bilateral microinjection of DADLE (10 nM in 50 nl) into the rVLM of the normotensive rats also did not influence BP. SBP, DBP and MBP were the same before and after microinjection of DADLE in the normotensive rats (143 ± 9, 79 ± 5 and 100 ± 6 mmHg). Heart rate was not affected by either δ-opioid receptor stimulation or antagonism.

**Microinjection sites.** Examination of the rat brain slices verified that each of the injections was located within the rVLM, with the exception of three ICI 174,864 injections that were found to be outside the rVLM of EA-treated hypertensive rats. None of the injections outside the rVLM altered blood pressure. Thus, microinjection sites observed to be 1.0 to 1.5 mm rostral to the obex, 0.5 to 1.0 mm from the ventral surface and 1.8 to 2.3 mm lateral to the midline, were considered to be within the rVLM, as noted in the atlas of Paxinos and Watson22 (Fig. 5).

**Discussion**

We have shown in a number of studies that EA inhibits sympathoexcitatory cardiovascular responses19–21,23. Preproenkephalin in the rVLM is increased following repetitive stimulation of the P5-6 acupoints and the underlying median nerves of normotensive rats24. However, we know little about mechanisms underlying EA’s influence in sustained hypertension. As such, we established a hypertensive model known to increase BP through its sympathoexcitation, to investigate the influence of EA in sustained hypertensive rats and to determine if opioids in
the brain stem underlie EA’s actions. The major new findings of this study are: (1) Repetitive EA reduces sustained hypertension, (2) Reduced BP in EA-treated rats persists for at least additional three days after end of EA treatment, providing evidence for a long-lasting hemodynamic action of EA, (3) EA increases the enkephalin gene expression in the rVLM for at least three days after eleven applications and (4) Blockade of δ-opioid receptors using ICI 174,864 or naloxone in the rVLM in EA-treated hypertensive rats partly reverses the EA response while stimulation of δ-opioid receptors using exogenous DADLE in the rVLM lowers BP in sham-EA hypertensive rats, directionally mimicking the EA effect in CIH. These data suggest that the δ-opioid system in the rVLM participates in the hypotensive action of EA in rats with sustained hypertension.

The sympathetic nervous system is essential for maintaining body temperature of cold-exposed rats. The responses of intact rats to cold exposure (4°C) include vasoconstriction, piloerection, shivering, adrenocorticotropin (ACTH) hypersecretion and increased mobilization of free fatty acids and glucose. Chemical sympathectomy blocks all of these responses except for ACTH hypersecretion. Animals subjected to chemical sympathectomy lose body heat rapidly and die in a few hours during cold exposure. However, the damaging actions of the chemical sympathectomy are reversed by administration of catecholamines. These studies suggest that increased sympathetic outflow contributes to development and maintenance of CIH. Norepinephrine in plasma increases after 24 hr of exposure to cold and remains elevated throughout the experiment, whereas the concentration of epinephrine in plasma increases early on but returns to control level after 19 days of exposure to cold. We also observed that BP remained elevated even when animals were placed in a neutral environment at room temperature for several hours during application of EA and experimental procedures. Compared

Figure 4. Involvement of rVLM δ-opioid receptors in EA modulation of blood pressure (BP) in rats with cold-induced hypertension (CIH). Panels A,B: δ-opioid receptor blockade in the rVLM reversed EA’s action in lowering BP in CIH rats. Panels C,D: decrease in BP after activation of δ-opioid receptors in the rVLM of CIH rats treated with sham-EA. Panels A,C show original BP tracings before and after bilateral microinjection of a δ-opioid receptor antagonist (ICI 174,864; 1 mM in 50 nl) and agonist (DADLE; 10 nM in 50 nl) into the rVLM. Arrows in Panels A,C indicate sequential bilateral microinjections into the rVLM. In Panels B,D, *indicates significant differences compared to before microinjection, *P < 0.05; **P < 0.01.
to the other forms of hypertension, such as spontaneous hypertension\(^3\), DOCA salt hypertension\(^6\) and renal hypertension\(^1\), CIH is presently the only "naturally-occurring" form of experimentally-induced hypertension that can be induced without surgical intervention, administration of excessive doses of drugs or hormones or genetic manipulation\(^1\). Also, as noted above, this form of hypertension has a large sympathoexcitatory component. As shown in our study and by others\(^13,24\), systolic, diastolic and mean BPs increase significantly after four weeks of exposure to mild cold (6°C) and are sustained at an elevated level for at least another six weeks in cold environments. Importantly, BP elevation is also observed in human subjects living in cold regions and during cold seasons\(^10,29\). Considering that EA inhibits sympathoexcitatory responses\(^19,20,23\), we predicted that EA would reduce the chronically elevated sympathoexcitatory state in the CIH model. Acupuncture does not influence thermosensation independently, i.e., more than controls\(^30–32\), although it may raise the pain threshold\(^32\), including thermal pain caused by excessive warm or cold\(^33,34\). Since mild cold (6°C) used in the present study was much less intense than the excessive cold that causes pain, it is unlikely that our observations of EAs action on BP were due to changes in cold threshold.

Stimulation of the ST36-37 (Zusanli-Shangjuxu) acupoints overlying the deep peroneal nerves has been widely used in treating several clinical conditions, including pain, spinal cord injury, nausea, gastrointestinal diseases, allergic rhinitis and cardiovascular conditions, such as hypertension\(^7,35\). We have shown that EA at ST36-37 or P5-6 similarly inhibit sympathoexcitatory reflex responses induced by gastric distension\(^26\) and gall bladder stimulation\(^29\). Both P5-6, overlying the median nerves, and ST36-37, overlying the deep peroneal nerves, inhibit sympathoexcitatory responses for prolonged periods of time and with similar intensities\(^16,37\). Although EA at ST36-37 and P5-6 appears to ameliorate BP in patients with mild to moderate hypertension for at least four weeks after terminating treatment\(^7\), the underlying mechanisms are unknown. We chose ST36-37 rather than P5-6 stimulation, since this approach was technically easier in conscious animals.

Our previous clinical study demonstrated that weekly EA therapy reduces elevated SBP and DBP, but not HR in patients with mild to moderate essential hypertension, and that EA had a stronger effect on elevations in SBP than DBP\(^7\). The efficacy of EA in lowering SBP may be important for patients with advancing age (≥60 years)\(^38\) since stress-related spikes in SBP places patients with hypertension at risk for stroke, developing peripheral artery disease, heart failure, as well as myocardial infarction and other cardiovascular diseases\(^38\). Thus, reduction in BP with EA has the potential to reduce the risk for these cardiovascular disorders. In the present study, we likewise found that EA reduced both SBP and DBP in animals with cold-induced hypertension. The present experimental study provides possible mechanisms by which EA reduces BP in patients with hypertension.

The rVLM serves an important role in cardiovascular sympathoexcitatory regulation\(^17,18\) and is an important site of EAs action in suppressing sympathoexcitation\(^6\). In this regard, we have shown that the opioid system in the rVLM including endorphin and enkephalins, that largely act through stimulation of \(\mu\)- and \(\delta\)-opioid receptors, respectively, contributes importantly to the sympathoinhibitory effects of EA on pressor reflex\(^19\). Anatomical studies have documented that enkephalins but not endorphin is synthesized in EA-activated neurons in the rVLM\(^19,40\). Endorphins are synthesized in the arcuate nucleus of the hypothalamus and are transported to the rVLM through long projections\(^39,40\).

Acupuncture’s long-lasting modulation of cardiovascular excitatory reflexes is mediated, in part, by opioids in the rVLM. In this regard, one 30 min session of EA in unanesthetized animals inhibits elevated BP for four or more hrs\(^41\). Under anesthesia, the inhibitory effect of 30 min of EA on premotor sympathoexcitatory cardiovascular rVLM neuronal responses lasts 60-90 min\(^19,21\). In the rVLM, opioids and GABA, but not nociceptin, participate in EA-related prolonged inhibition of sympathoexcitatory cardiovascular responses\(^22\). Furthermore, mRNA expression of preproenkephalin in the rVLM is increased at 90 min after 30 min of EA and 24 hr after two repetitive applications of EA in anesthetized\(^42\) and conscious normotensive rats\(^23\), suggesting that molecular
changes of enkephalin gene expression in the rVLM may contribute to EAs immediate actions. Despite these
prior observations, it has been unclear if increases in preproenkephalin in the rVLM in response to repetitive EA
over a five week period contribute to the BP long-lasting lowering effect of EA in rats with sustained hypertension.
In this regard, the current study is the first to demonstrate that repeated application of EA reduces sustained hyperten
sion induced by cold stress for at least three days after the course of EA treatment. This prolonged action is
associated with elevated preproenkephalin mRNA expression levels in the rVLM. The observation that pre-
proenkephalin is increased in the rVLM after repeated EA in the hypertensive rats, suggests that EA-associated
increases in expression of preproenkephalin enhances the availability of rVLM enkephalin to, in turn, decrease elevated neuronal activity, sympathetic outflow and ultimately hypertension. Our previous study documents that the protein (met-enkephalin) rises in parallel with its mRNA precursor, preproenkephalin, following repetitive EA44. These data in aggregate therefore suggest that molecular changes of enkephalin gene expression are related to EAs long-lasting actions on elevated BP in hypertensive models.

Microinjection of a δ-opioid receptor agonist into the rVLM, like EA, decreased elevated BP in CIH rats, while
ICI 174,864, the δ-opioid receptor antagonist partly reverses the EA effect. Thus, increased opioid synthesis in the
rVLM associated with EA participates in the long-lasting inhibitory action of EA in CIH.

Microinjection of ICI 174,864, the δ-opioid receptor antagonist, or DARL, the δ-opioid agonist, into the
rVLM of normotensive rats did not influence blood pressure. This result is consistent with our and other previous
studies and indicates that there is no tonic action resulting from δ-opioid receptor stimulation18,20,23,24,43–45. Thus
it appears that under normal physiological conditions endogenous opioids in the rVLM do not participate in BP
regulation. Furthermore, endogenous opioid release induced by EA takes place without influencing baseline BP. It is
likely that in normotensive animals, sympathetic excitatory cardiovascular neurons in rVLM maintain a low level of
activation and thus there is little for exogenous opioids to act upon. On the other hand, when rVLM neurons are
activated during reflex responses18,20,23,24,43–45 or in the setting of hypertension45, exogenous opioids administered
through microinjection or increased endogenous opioids in the rVLM in response to EA decrease pressor reflex
responses and attenuate hypertension46,47. As such, increased expression of opioid receptors during the
development of hypertension also contributes to an enhanced opioid system in the rVLM.

Alpha-chloralose frequently is used in cardiovascular experimentation19,23,44,48 since it preserves cardiovascular
reflex function better than other anesthetics like barbiturates49,50. The actions of enkephalin on BP under
α-chloralose anesthesia are similar to those observed in conscious preparations51. Under urethane anesthesia,
100 nl of 0.2 mM endorphin-2 microinjected into the rVLM of normotensive Wistar rats has been shown to
decrease BP and heart rate52. Also, in α-chloralose and urethane anesthetized rats, microinjection of 100 nl of
3 nM D-Ala²-N-Me-Phe³-Gly⁵-ol-Enkephalin acetate salt (DAGO), a μ-opioid agonist or 10 nM naloxone into
the rVLM of normotensive Sprague Dawley rats respectively decreases or increases arterial pressure and heart rate.
The reasons for these discrepant observations are uncertain but might be related to the different anesthetics and/or
microinjection volume.

Hypertension affects about one third of the adult population of the world. It is also the leading chronic risk
factor for mortality7. Acupuncture potentially presents a viable non-pharmacological treatment option. However,
due to limitations of the previous clinical studies, including small sample sizes, methodological and design het-
erogeneity, and differences in BP related end-points, the actions of acupuncture in hypertension remains con-
troversial17. Using a clinically relevant model of hypertension, the current study demonstrates that repetitive EA
attenuates hypertension by about 40% through a δ-opioid mechanism, occurring at least in part in the rVLM.
Repetitive EA evokes a long-lasting response, suggesting that this therapy may be suitable for treating clinical
hypertension.

In conclusion, repetitive EA attenuates CIH. The BP lowering action of EA lasts at least three days after ces-
sation of EA. The inhibitory effect of EA is related to the increases in enkephalin levels in the rVLM. The present
study suggests that acupuncture may be a viable non-pharmacological option to treat chronic hypertension.

Methods
The study conformed to the American Physiological Society's Guiding Principles for Research Involving Animals
and Human Beings. Experimental protocols were approved by the Institutional Animal Care and Use Committee
of the University of California at Irvine. A total of 54 male Sprague Dawley rats (150 g) were purchased from
Charles River Laboratories International, Inc.). Thirty-six rats were placed into a cold room (6 °C) while 18 rats
maintained in room temperature (25 °C) for 11 weeks. They all were provided with standard rodent chow and
tap water ad libitum and housed in 12-12 hr light-dark cycle. Figure 1 displays the timeline of the experiments.

Blood pressures and heart rates in conscious rats were measured non-invasively with a volume pressure
recording sensor and an occlusion tail-cuff (CODA System, Kent Scientific). Digital values were recorded. To
evaluate BP, each animal was placed in a restrainer, and the cuff was placed on the tail and inflated and released
several times to condition the rat to the procedure. After stabilization, BP was measured five times to acquire an
average BP. BPs and HRs in all rats were evaluated weekly (Fig. 1).

Experimental protocols. All rats were gently restrained in slings for 30 min twice weekly for six weeks to
accommodate them to manipulation and to prevent stress associated with this procedure. Each rat was wrapped
with a cloth around the body, excluding the four limbs. This method allowed manipulation of the limbs and
application of EA at ST36–37 acupoints both safely and effectively in the immobilized conscious rats. The ST36–37
acupoints (overlying deep peroneal nerve below the knee on the hind limb) are analogous to those in humans53.
Acupuncture needles (0.16 mm diameter) were inserted into the ST36–37 acupoints briefly (~ 10 s) to allow them
to become accustomed to handling and the acupuncture procedure. The rat maintained at room temperature was
handled without needle insertion.
After six weeks the rats housed in cold room demonstrated elevated SBP compared to rats maintained at room temperature. The CIH rats were divided randomly into EA, sham-EA or untreated hypertensive groups.

**EA-treated group.** Acupuncture needles were inserted bilaterally in ST36-37 and stimulated with a constant current stimulator connected to a stimulus isolation unit (Grass, model S88) to provide EA (2 Hz, 0.1 – 0.4 mA, 0.5 ms duration) for 30 min\(^{21,24,42}\), twice weekly for five weeks (Fig. 1). Correct positioning of the needles at ST36-37 was confirmed by observing slight repetitive paw twitches, indicating stimulation of motor fibers in the mixed nerve bundle comprising the deep peroneal nerve\(^{44}\). Of note, motor nerve stimulation does not participate in the EA-cardiovascular response\(^{54}\).

**Sham-EA group.** The CIH rats were treated the same as that in the EA group with the exception that needles were inserted but not electrically stimulated.

**Hypertensive control group.** These animals were restrained 30 min twice a week but were not subjected to insertion of acupuncture needles (Fig. 1).

**Normotensive control group.** Rats were maintained at room temperature for 11 weeks with 30 min restraint twice weekly (Fig. 1). No acupuncture was applied to this group.

### Expression of preproenkephalin mRNA in the rVLM

**Tissue collection from rVLM.** The rVLM tissue was collected 72 hr following five-weeks of treatment with EA (n = 9) and sham-EA (n = 6), as well as from six hypertensive and six normotensive rats (Fig. 1). All rats were anesthetized with ketamine/xylazine (75–100/5–10 mg/kg, i.p.), then decapitated. Brain tissue was removed and stored in RNA Later solution (Ambion). The rVLM tissue was obtained by positioning a 20-gauge needle stub 0.5 mm lateral to the lateral edge of the pyramid tract next to the caudal edge of the trapezoid body from ventral surface of the brain stem\(^{24,42}\). The needle stub was inserted perpendicularly 0.5 mm towards dorsally. The first 0.2 mm of harvested tissue from the ventral surface was excluded because it was located outside the rVLM, and the remaining tissue was confirmed to be in the rVLM\(^{22,24,42}\). Bilateral rVLM tissue samples were collected from each rat.

**RNA isolation and real time PCR.** Tissue samples were homogenized in a glass tissue grinder (DUALL 20, Kontes Glass Co.) using 800μl Trizol reagent (Life Technologies). RNA was dissolved in 10μl nuclease-free water, and the concentration and purity of RNA was determined spectrophotometrically using a Nanodrop ND-1000 spectrophotometer. One hundred ng of total RNA were transcribed using SuperScript II RT and a mixture of oligo (dT) (100 ng/reaction) and random primers (200 ng/reaction; all from Invitrogen). Real-time quantitative PCR was performed with an Opticon 4 using the SYBR green real-time master mix (Bio-Rad). The following primer sequences were used: preproenkephalin, forward (5′-tgc ctt ctt tca aaa tct gg-3′); reverse (5′-ggg gta aag ctc atc cat ct-3′); 18s, forward (5′-tgc ctt ctt tca aaa tct gg-3′); reverse (5′-acg cta tca tca tca-3′). The sizes of the PCR products for preproenkephalin was 191 bp and for the 18s housekeeping gene 185 bp. The PCR program was 95 °C 10 min, (95 °C 10 s, 59 °C 45 s) x 50 cycles followed by the melting curve analysis (55–90 °C) to verify the PCR products for preproenkephalin was 191 bp and for the 18s housekeeping gene 185 bp. The PCR program was 95 °C 10 min, (95 °C 10 s, 59 °C 45 s) x 50 cycles followed by the melting curve analysis (55–90 °C) to verify product specificity. For each sample, the copy number of both preproenkephalin and the 18s housekeeping gene was extrapolated from their respective standard curves. The value of preproenkephalin mRNA expression was normalized by the number of 18s copies and expressed in arbitrary units. Reproducibility of results was determined by performing triplicate measurements of each cDNA aliquot.

### Role of rVLM δ-opioid receptors in EA-modulated hypertension

**Drugs.** Naloxone (1 mM) was purchased from Sigma-Aldrich. ICI 174,864 (1 mM)\(^{55}\) and DADLE (10 nM)\(^{56}\) were δ-opioid receptor antagonist and agonist, respectively. Both of them were purchased from Tocris. ICI 174,864 (5 mg) was dissolved in 100μl DMSO and then diluted to a concentration of 1 mM with 0.9% normal saline. Naloxone and DADLE were dissolved in the normal saline.

**Surgical preparation.** Anesthesia was induced with ketamine (100 mg/kg, i.m.) and α-chloralose (50–60 mg/kg, i.v.). Additional doses of α-chloralose (25–30 mg/kg, i.v.) were administered to maintain an adequate level of anesthesia. A femoral vein and artery were cannulated for administration of fluids and monitoring BP, respectively. HR was derived from the pulsatile BP signal. The trachea was intubated to provide artificial ventilation using a respirator (model 661, Harvard Apparatus). Arterial blood gases were measured with a blood gas analyzer (ABL5, Radiometer America) and maintained within the normal physiological range (PO2 >100 mmHg, PCO2 30–40 mmHg and arterial pH 7.35–7.4). Body temperature was maintained between 36 and 38 °C with a heating pad.

**Microinjections in the rVLM.** Animals were placed in a stereotaxic head frame. A partial craniotomy was performed to expose the dorsal medulla. Two CMA microdialysis probes (14 mm long with 0.24 mm of tip diameter) used for microinjections were modified by removing microdialysis membrane\(^{43}\). They were inserted bilaterally into the rVLM, 1.8–2.3 mm lateral from the midline, 1–1.5 mm rostral to the obex, and advanced 3.0–3.3 mm\(^{21,22,27}\). The probes were connected to 25μl Hamilton syringes fitted in UMP3 microsyringe injectors (WPI). The microinjections were conducted sequentially into bilateral rVLM.

**Opioid antagonism in EA CIH rats.** The naloxone (n = 2) or ICI 174,864 (n = 4; 1 mM in 50 nl) were bilaterally microinjected into the rVLM of CIH rats subjected to eleven repeated applications of EA. This dose of ICI 174,864 in the rVLM attenuates EA modulation of the BP response\(^{49}\). Vehicle controls were examined in four rats.
Delta-opioid receptor stimulation in sham-EA CIH rats. The BP response to rVLM stimulation with DADLE (10 nM in 50 nl) was examined in six sham-EA rats. This dose of DADLE injected into the rVLM markedly lowers BP in acute and sustained hypertension29,46. In four of these rats, the normal saline was also microinjected into the rVLM two hrs either before or after administration of DADLE.

Activation and inhibition of delta-opioid receptor in normotensive rats. DADLE or ICI 174,864 was microinjected bilaterally into the rVLM in six normotensive rats.

Histological confirmation of microinjection sites. We injected 5% Chicago Sky Blue dye (100 nl) at the end of each experiment to identify probe placement. After euthanasia, the medulla oblongata was removed and submerged in 4% paraformaldehyde. Coronal sections (40 μm) were cut with a cryostat microtome (Leica). Using the rat’s atlas22 as a guide, sites of microinjections in the medulla were plotted with Corel Presentation software on reconstructed coronal sections24,43.

Statistical analysis

Data are presented as means ± SEM. A one-way repeated measure of ANOVA followed post hoc by the Student–Newman Keuls test was used to compare changes in BPs and HRs over time as well as before and after drug microinjection in each group. The difference of BP and HR between two groups was compared using a two-way repeated measure ANOVA followed the Holm–Sidak t-test. Comparisons of preproenkephalin mRNA level between two groups were analyzed using the Student’s t-test. Statistical analyses were performed with Sigma Plot (Jandel Scientific). The 0.05 probability level was used to detect significant differences.

References

1. Kearney, P. M. et al. Global burden of hypertension: analysis of worldwide data. Lancet 365, 217–223 (2005).
2. Chobanian, A. V. et al. Seventh report of the Joint National Committee on Prevention, Detection, Evaluation, and Treatment of High Blood Pressure. Hypertension 42, 1206–1252 (2003).
3. Macklin, E. A. et al. Stop Hypertension with the Acupuncture Research Program (SHARP): results of a randomized, controlled clinical trial. Hypertension 48, 838–845 (2006).
4. Yin, C. et al. Acupuncture, a promising adjunctive therapy for essential hypertension: a double-blind, randomized, controlled trial. NeuroRes 29 Suppl 1, S98–103 (2007).
5. Brook, R. D. et al. Beyond medications and diet: alternative approaches to lowering blood pressure: a scientific statement from the american heart association. Hypertension 61, 1360–1383 (2013).
6. Longhurst, J. C. & Tjen-A-Looi, S. C. Acupuncture regulation of blood pressure: two decades of research. Int. Rev. Neurobiol. 111, 257–271 (2013).
7. Li, P. et al. Long-lasting reduction of blood pressure by electroacupuncture in patients with hypertension: randomized controlled trial. Medical Acupuncture 27, 253–266 (2015).
8. Carlsson, C. P. & Sjöland, B. H. Acupuncture for chronic low back pain: a randomized placebo-controlled study with long-term follow-up. Clin J Pain 17, 296–305 (2001).
9. Guerra de Hoyos, J. A. et al. Randomised trial of long term effect of acupuncture for shoulder pain. Pain 112, 289–298 (2004).
10. Hata, T. et al. The seasonal variation of blood pressure in patients with essential hypertension. Clin Exp Hypertens A 4, 341–354 (1982).
11. Cold exposure and winter mortality from ischaemic heart disease, cerebrovascular disease, respiratory disease, and all causes in warm and cold regions of Europe. The EuroWinter Group. Lancet 349, 1341–1346 (1997).
12. Yang, L. et al. Outdoor temperature, blood pressure, and cardiovascular disease mortality among 23 000 individuals with diagnosed cardiovascular diseases from China. Eur Heart J 36, 1178–1185 (2015).
13. Fregly, M. I., Kitka, D. C., Threette, R. M., Torres, J. L. & Barney, C. C. Development of hypertension in rats during chronic exposure to cold. J Appl Physiol 66, 741–749 (1989).
14. Papane, P. E., Wood, C. E. & Fregly, M. J. Role of the sympathetic nervous system in cold-induced hypertension in rats. J Appl Physiol 71, 300–306 (1991).
15. Maickel, R. P., Matsusak, N., Stern, D. N. & Brodie, B. B. The sympathetic nervous system as a homeostatic mechanism. I. Absolute need for sympathetic nervous function in body temperature maintenance of cold-exposed rats. J Pharmacol Exp Ther 157, 103–110 (1967).
16. Sun, Z. J. & Zhang, Z. E. Historic perspectives and recent advances in major animal models of hypertension. Acta Pharmacol Sin. 26, 295–301 (2005).
17. Kumagai, H. et al. Importance of rostral ventrolateral medulla neurons in determining efferent sympathetic nerve activity and blood pressure. Hypertens. Res. 35, 132–141 (2012).
18. Goyenet, P. G. The sympathetic control of blood pressure. Nat. Rev. Neurosci. 7, 335–346 (2006).
19. Li, P., Tjen-A-Looi, S. C. & Longhurst, J. C. Rostral ventrolateral medullary opioid receptor subtypes in the inhibitory effect of electroacupuncture on reflex autonomic response in cats. Autonomic Neuroscience 89, 38–47 (2001).
20. Tjen-A-Looi, S. C., Li, P. & Longhurst, J. C. Prolonged inhibition of rostral ventral lateral medullary premotor sympathetic neuron by electroacupuncture in cats. Auton Neurosci.: Basic and Clinical 106, 119–131 (2003).
21. Zhou, W., Tjen-A-Looi, S. & Longhurst, J. C. Brain stem mechanisms underlying acupuncture modality-related modulation of cardiovascular responses in rats. J Appl Physiol, 99, 851–860 (2005).
22. Paxinos, G. & Watson, C. The rat brain in stereotaxic coordinates. Academic Press (2009).
23. Tjen-A-Looi, S. C., Li, P. & Longhurst, J. C. Role of medullary GABA, opioids, and nociceptin in prolonged inhibition of cardiovascular sympathoexcitatory reflexes during electroacupuncture in cats. Am J Physiol 293, H3627–H3635 (2007).
24. Li, M., Tjen-A-Looi, S. C., Guo, Z. L. & Longhurst, J. C. Repetitive electroacupuncture causes prolonged increased met-enkephalin expression in the rVLM of conscious rats. Auton. Neurosci 170, 30–35 (2012).
25. Judy, W. V. & Farrell, S. K. Arterial baroreceptor reflex control of sympathetic nerve activity in the spontaneously hypertensive rat. Hypertension 1, 605–614 (1979).
26. Nakamura, Y. et al. Central attenuation of aortic baroreflex inhibition in prehypertensive DOCA-salt-loaded rats. Hypertension 12, 259–266 (1988).
27. Guo, G. B., Thames, M. D. & Abbood, F. M. Arterial baroreflexes in renal hypertensive rabbits. Selectivity and redundancy of baroreceptor influence on heart rate, vascular resistance, and lumbar sympathetic nerve activity. Circ Res 53, 223–234 (1983).
28. Wang, X. et al. AAV-based RNAi silencing of NADPH oxidase gp91(phox) attenuates cold-induced cardiovascular dysfunction. Hum Gene Ther 23, 1016–1026 (2012).
29. Brennan, P. J., Greenberg, G., Miall, W. E. & Thompson, S. G. Seasonal variation in arterial blood pressure. Br Med J (Clin Res Ed) 285, 919–923 (1982).
30. Downs, N. M., Kirk, K. & MacSweeney, A. The effect of real and sham acupuncture on thermal sensation and thermal pain thresholds. Arch. Phys. Med. Rehabil. 86, 1252–1257 (2005).
31. Amand, M., Nguyen-Huu, F. & Balestra, C. Acupuncture effect on thermal tolerance and electrical pain threshold: a randomised controlled trial. *Acupunct. Med.* **29**, 47–50 (2011).
32. Lundeborg, T., Eriksson, S., Lundeborg, S. & Thomas, M. Acupuncture and sensory thresholds. *Am. J. Chin. Med.* **17**, 99–110 (1989).
33. Leung, A., Khadi, B., Duann, J. R., Cho, Z. H. & Yaksh, T. The effect of Ting point (tendomuscular meridians) electroacupuncture on thermal pain: a model for studying the neuronal mechanism of acupuncture analgesia. *J. Altern. Complement Med.* **11**, 653–661 (2005).
34. Wang, S. M., Lin, E. C., Maranets, I. & Kain, Z. N. The impact of asynchronous electroacupuncture stimulation duration on cold thermal pain threshold. *Anesth. Analg.* **109**, 932–935 (2009).
35. Stux, G. & Pomeranz, B. *Basics of acupuncture*. Springer-Verlag, Berlin (1998).
36. Zhou, W., Fu, L.-W., Tjen-A-Looi, S. C., Li, P. & Longhurst, J. C. Affenter mechanisms underlying stimulation modality-related modulation of acupuncture-related cardiovascular responses. *J Appl Physiol* **98**, 872–880 (2005).
37. Tjen-A-Looi, S. C., Li, P. & Longhurst, J. C. Medullary substrate and differential cardiovascular response during stimulation of specific acupoints. *Am J Physiol* **287**, R852–R862 (2004).
38. Rapsomaniki, E. et al. Blood pressure and incidence of twelve cardiovascular diseases: lifetime risks, healthy life-years lost, and age-specific associations in 1.25 million people. *Lancet* **383**, 1899–1911 (2014).
39. Guo, Z.-L., Mozazami, A. R. & Longhurst, J. C. Electroacupuncture induces c-Fos expression in the rostral ventrolateral medulla and periaqueductal gray in cats: relation to opioid containing neurons. *Brain Res.* **1030**, 103–115 (2004).
40. Li, P., Tjen-A-Looi, S. C., Guo, Z.-L., Fu, L.-W. & Longhurst, J. C. Long-loop pathways in cardiovascular electroacupuncture responses. *J Appl. Physiol.* **106**, 620–630 (2009).
41. Yao, T., Andersson, S. & Thoren, P. Long-lasting cardiovascular depression induced by acupuncture-like stimulation of the sciatic nerve in unanaesthetized spontaneously hypertensive rats. *Brain Res.* **240**, 77–85 (1982).
42. Li, M., Tjen-A-Looi, S. C. & Longhurst, J. C. Electroacupuncture enhances preproenkephalin mRNA expression in rostral ventrolateral medulla of rats. *Neurosci Lett* **477**, 61–65 (2010).
43. Li, M., Tjen-A-Looi, S. C., Guo, Z.-L. & Longhurst, J. C. Electroacupuncture modulation of reflex hypertension in rats: role of cholecystokinin octapeptide. *Am J Physiol 305*, R404–R413 (2013).
44. Chao, D. M. et al. Naltrexone reverses inhibitory effect of electroacupuncture on sympathetic cardiovascular reflex responses. *Am. J. Physiol.* **276**, H2127–H2134 (1999).
45. Zamir, N., Simantov, R. & Segal, M. Pain sensitivity and opioid activity in genetically and experimentally hypertensive rats. *Brain Res* **184**, 299–310 (1980).
46. Sun, S. Y., Liu, Z., Li, P. & Ingenito, A. J. Central effects of opioid agonists and naltrexone on blood pressure and heart rate in normotensive and hypertensive rats. *Gen Pharmacol* **27**, 1187–1194 (1996).
47. Li, P., Zhe, D. M., Kao, K. M., Lin, Q. & Sun, S. Y. Role of acetylcholine, corticoids and opioids in the rostral ventrolateral medulla in stress-induced hypertensive rats. *Biol Signals* **4**, 124–132 (1995).
48. Cox, R. H. Influence of chloralose anesthesia on cardiovascular function in trained dogs. *Am J Physiol* **223**, 660–667 (1972).
49. Shebata, R., Fowler, N. & HURLBURT, O. Hemodynamics studies of dogs under pentobarbitol and morphine chloralose anesthesia. *J Surg Res* **3**, 263–267 (1963).
50. Priano, L. L., Traber, D. L. & Wilson, R. D. Barbiturate anesthesia: an abnormal physiologic stimulation. *J Pharmacol Exp Ther* **165**, 126–135 (1969).
51. Sander, G. E., Lowe, R. F. & Giles, T. D. The effects of barbiturates upon the hemodynamic responses to intravenous methionine-enkephalin in dogs: modulation by the GABA complex. *Peptides* **7**, 259–265 (1986).
52. Kasamatsu, K. & Sapru, H. N. Attenuation of aortic baroreflex responses by microinjections of endorphin-2 into the rostral ventrolateral medullary pressure area of the rat. *Am J Physiol Regul Integr Comp Physiol* **289**, R59–R67 (2005).
53. Hua, X. B. Acupuncture manual for small animals experimental acupuncture. Shanghai Science and Technology Publisher, Shanghai (1994).
54. Li, P., Tjen-A-Looi, S. C. & Longhurst, J. C. Excitatory projections from arcuate nucleus to ventrolateral periaqueductal gray in unanaesthetized spontaneously hypertensive rats. *Brain Res* **924**, R59–R67 (2001).
55. Cotton, R., Giles, M. G., Miller, L., Shaw, J. S. & Timms, D. ICI 174864: a highly selective antagonist for the opioid delta-receptor. *Eur J Pharmacol* **97**, 331–332 (1984).
56. Suh, H. H. & Tseng, L. F. Different types of opioid receptors mediating analgesia induced by morphine, DAMGO, DPDPE, DADLE and beta-endorphin in mice. *Naunyn Schmiedebergs Arch Pharmacol* **342**, 67–71 (1990).
57. Tjen-A-Looi, S. C., Guo, Z.-L., Li, M. & Longhurst, J. C. Medullary GABAergic mechanisms contribute to electroacupuncture modulation of cardiovascular depressor responses during gastric distention in rats. *Am J Physiol 304*, R321–R332 (2013).

**Acknowledgements**

We are sincerely thankful to Keun Kim, Ai-thuann Nguyen, Jonathan Chou, Ye Na Choi, Shushanna Galstyan and Gohar Khechumyan for their technical assistance. This study was supported by NIH grants, HL-072125 (S.C. Tjen-A-Looi and J.C. Longhurst) and AT009347 (Z.-L. Guo and J.C. Longhurst), AHA 10POST4190125 (M. Li) and Department of Medicine Chair Research Award at University of California, Irvine (Z.-L. Guo).

**Author Contributions**

M.L. and Z.-L. G contributed to data collection, data analyses and writing of manuscript. S.C.T.-A.L. contributed to conceptualization of experiments and data analyses. J.C.L. contributed to conceptualization of experiments and review of data. All authors reviewed and approved the manuscript.

**Additional Information**

**Competing financial interests:** The authors declare no competing financial interests.

**How to cite this article:** Li, M. *et al.* Repetitive Electroacupuncture Attenuates Cold-Induced Hypertension through Enkephalin in the Rostral Ventral Lateral Medulla. *Sci. Rep.* **6**, 35791; doi: 10.1038/srep35791 (2016).

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