Involvement of the lateral septal area in the expression of fear conditioning to context

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Considering the evidence that the lateral septal area (LSA) modulates defensive responses, the aim of the present study is to verify if this structure is also involved in contextual fear conditioning responses. Neurotransmission in the LSA was reversibly inhibited by bilateral microinjections of cobalt chloride (CoCl₂, 1 mM) 10 min before or after conditioning or 10 min before re-exposure to the aversively conditioned chamber. Only those animals that received CoCl₂ before re-exposure showed a decrease in both cardiovascular and behavioral conditioned responses. These results suggest that the LSA participates in the expression, but not acquisition or consolidation, of contextual fear conditioning.

The lateral septal area (LSA) has been related to the modulation of several cognitive and emotional processes including learning, memory, anxiety, and regulation of autonomic responses (Covian 1966; Paxinos 1995; Sheehan et al. 2004; Scopinho et al. 2006, 2007). This structure is activated during aversive situations (Pezone et al. 1992; Duncan et al. 1993; Beck and Fibiger 1995; Kubo et al. 2002) and sends projections to brain regions involved in the behavioral and cardiovascular responses to aversive stimuli (LeDoux et al. 1988; Resstel et al. 2006a, 2008a,c; Tavares and Correa 2006; Tavares et al. 2009). There is evidence that the LSA modulates the autonomic responses to stress and emotional threat situations (Kubo et al. 2002). Also, the anxiolytic-like effect evoked by systemic administration of diazepam in rats submitted to fear conditioning to context is associated with a decrease in LSA neuronal activity (Beck and Fibiger 1995). Taken together, these data support a possible regulatory role of the LSA on behavioral and cardiovascular responses associated with aversive situations such as fear conditioning.

Conditioned fear to context is evoked by re-exposing an animal to an environment (context) that has been previously paired with an aversive or unpleasant stimulus (Blanchard and Blanchard 1969; Fanselow 1980, 2000; Resstel et al. 2006b). This re-exposure causes freezing immobility and increases in mean arterial pressure (MAP) and heart rate (HR) (Fanselow 1980; LeDoux et al. 1988; Resstel et al. 2008a,b). These responses are modulated by structures connected with the LSA, such as the medial prefrontal cortex, amygdala, and bed nucleus of the stria terminalis (Swanson and Cowan 1977; Risold and Swanson 1997; Kuniecki et al. 2002; Vertes 2004; Resstel et al. 2008a,b). Therefore, the LSA could be part of the brain circuitry involved in contextual fear conditioning. However, the precise role of the LSA in contextual fear conditioning is still not completely understood, with contradictory evidence found in the literature. For example, freezing behavior elicited by an aversively conditioned context was reported to be potentiated in rats with LSA lesion (Sparks and LeDoux 1995), whereas a recent study showed that LSA reversible inhibition by lidocaine before the conditioning session reduces freezing (Calandreau et al. 2007). Since the former study employed irreversible LSA lesions that could have, in addition to acquisition, also influenced consolidation, retrieval, or expression of the aversive memory, differences in the experimental protocols could be responsible for the contradictory results.

Therefore, the aim of the present study was to investigate the role of the LSA in the acquisition, consolidation, and retrieval of contextual fear memory by measuring the behavioral and cardiovascular responses evoked by re-exposure to an aversively conditioned context. To accomplish that, the nonselective neurotransmitter blocker cobalt chloride (CoCl₂) (Hagiwara and Byerly 1981; Lomber 1999) was microinjected into the LSA at the above-mentioned specific phases of the experimental procedure.

Male Wistar rats weighing 230–250 g were used. The animals were kept in the animal care unit of the Department of Pharmacology, School of Medicine of Ribeirão Preto, University of São Paulo, Brazil. The rats were housed individually in plastic cages with free access to food and water under a 12-h light/dark cycle (lights on at 06.30 h). The institution’s Animal Ethics Committee approved the housing conditions and experimental protocols.

Seven days before the experiment, the rats were anesthetized with 2,2,2-tribromoethanol (250 g/kg intraperitoneally; Sigma). After scalp anesthesia with 2% lidocaine, the skull was surgically exposed, and stainless steel guide cannulae (0.55 mm) were implanted bilaterally into the LSA using a stereotaxic apparatus (Stoeling), as described by Scopinho et al. (2007).

Preconditioning, conditioning, and testing were carried out in a 25 × 22 × 22-cm footshock box. The box had a grid floor composed of 18 stainless steel rods (2 mm in diameter), spaced 1.5 cm apart and wired to a shock generator (Automatic Reflex Conditioner, model 8572; Ugo Basile). The experimental box was cleaned with 70% ethanol before and after use. Preconditioning started 1 wk after guide cannula implantation and consisted of one 10-min-long pre-exposure (habituation) in the footshock box. In the conditioning shock session, performed 24 h after the habituation session, animals were divided into two experimental groups: nonconditioned and conditioned groups. The conditioned group was re-exposed to the footshock chamber, and, after 3 min of habituation, the animals were submitted to a...
At the end of the experiments the rats were sacrificed under deep anesthesia to injection site determination as described by Scopinho et al. (2007). A representative photomicrograph and a diagrammatic representation indicating the injection sites in the LSA can be seen in Figure 1.

There was a significant interaction between condition, group, and treatment ($F_{(2,69)} = 13.7, P < 0.001$). Exposure to the aversive context increased freezing in vehicle-treated conditioned animals compared to the nonconditioned groups. Intra-LSA injection of CoCl$_2$ failed to change this effect when administered immediately before or after the conditioning session. However, this treatment induced a significant decrease in freezing when administered before the test session (Bonferroni's test, $P < 0.05$; Fig. 2). CoCl$_2$ treatment did not change motor activity (number of crossings, $F_{(1,175)} = 1.5, P > 0.05$; rears, $F_{(1,177)} = 1.1, P > 0.05$) in nonconditioned animals. Pre-exposure to the conditioning chamber did not change the effects of CoCl$_2$ on the expression of contextual fear conditioning (data not shown).

No significant difference was found in baseline values of MAP and HR of conditioned and nonconditioned groups of animals (MAP: $F_{(1,155)} = 0.75, P > 0.05$; HR: $F_{(1,157)} = 1.7, P > 0.05$). Similar to the behavioral responses, there was a significant interaction between condition, group, treatment, and time (MAP: $F_{(45,69)} = 2.5, P < 0.05$; HR: $F_{(45,69)} = 2.7, P < 0.05$). The administration of CoCl$_2$ had no effect on cardiovascular responses observed during the chamber re-exposition in nonconditioned animals before conditioning (MAP: $F_{(1,135)} = 1.6, P > 0.05$; HR: $F_{(1,135)} = 0.14, P > 0.05$), after conditioning (MAP: $F_{(1,120)} = 0.66, P > 0.05$; HR: $F_{(1,120)} = 0.03, P > 0.05$), or before the test (MAP: $F_{(1,135)} = 0.2, P > 0.05$; HR: $F_{(1,135)} = 1, P > 0.05$; Fig. 3). Compared with the vehicle-treated group, injection of CoCl$_2$ in conditioned animals before or after conditioning did not change the increases in MAP ($F_{(1,150)} = 0.2, P > 0.05$ and $F_{(1,135)} = 1, P > 0.05$, respectively) and HR ($F_{(1,150)} = 0.2, P > 0.05$ and $F_{(1,135)} = 1, P > 0.05$, respectively) (Fig. 3). When administered before the test, however, CoCl$_2$ attenuated the increases in both MAP and HR induced by re-exposure to the context (MAP: $F_{(1,180)} = 137, P < 0.001$; HR: $F_{(1,180)} = 89, P < 0.001$) (Fig. 3).

Acquisition of contextual conditioned fear relates to the initial learning process that occurs when an aversive unconditioned stimulus (e.g., footshock) is associated with a specific context in the training session. This is followed by a memory consolidation phase. Subsequent re-exposure to the aversive context induces a retrieval of the conditioned memory, reflected by the expression of the conditioned fear responses (Fanselow 2000). Acute neurotransmission inhibition in the LSA by local injection of CoCl$_2$ before the test session reduced both freezing and cardiovascular responses observed during chamber re-exposure of the conditioned group. However, when CoCl$_2$ was injected 10 min before conditioning.

### Figure 1

**A** Photomicrograph of a coronal brain section showing bilateral microinjection sites in the lateral septal area (LSA). **B** Diagrammatic representation based on the rat brain atlas of Paxinos and Watson (1997) indicating injections sites of vehicle or CoCl$_2$ inside (●) or outside (○) the lateral septal area (LSA). (IA) Interaural; (cc) corpus callosum; (LV) lateral ventricle.

### Figure 2

Effects of bilateral microinjection of 100 nl of vehicle or 1 mmol of CoCl$_2$ performed at three different treatment periods on the percentage of time spent in freezing behavior in nonconditioned ($n = 5–6$) and conditioned animals ($n = 5–7$). Columns represent the means, and bars the SEM; (*) $P < 0.05$ compared to vehicle nonconditioned group; (#) $P < 0.05$ compared to vehicle conditioned group, Bonferroni's post-hoc test.
before or 10 min after the conditioning session, no interference in the expression of fear conditioning was found. These data suggest that synaptic neurotransmission in the LSA is involved in retrieval of the aversive context memory, but it is not implicated in acquisition or consolidation of aversive context memory. Because the treatment had no effect on nonconditioned rats, the results also indicate that the observed effects depend on a specific interaction with the fear conditioned response rather than nonspecific drug effects.

Irreversible lesions of the LSA have yielded contradictory results in animals exposed to aversive situations (Pesold and Treit 1992; Sparks and LeDoux 1995; Menard and Treit 1996; Sheehan et al. 2004). In contrast with our results, Sparks and LeDoux (1995) described that LSA electrolytic lesion performed 14 d before the conditioning session increased conditioned freezing behavior to context. However, in addition to including fibers of passage, the electrolytic lesions in their experiment are larger than ours, involving most parts of the septal area. Moreover, it

Figure 3. Time course of the effects of bilateral microinjection of 100 nL of vehicle or 1 nmol of CoCl₂ performed at three different treatment periods on mean arterial pressure (Δ MAP) and heart rate (Δ HR) in (top panel) nonconditioned (n = 5–6) and (bottom panel) conditioned animals (n = 5–7). Arrows indicate the start of the chamber re-exposure. Symbols represent the means, and bars the SEM. (*) P < 0.05, Bonferroni’s post-hoc test.
has been shown that the emotional changes that follow brain lesions may vary as a function of time (Rangel et al. 2003). Social interaction, for example, increased 2 wk after ventral medial prefrontal cortex (vMPFC) lesions, but decreased 5 wk later, suggesting the involvement of secondary plastic changes (Rangel et al. 2003). In the present study, we overcame these problems by inducing an acute and reversible LSA inhibition with CoCl2 (Scopinho et al. 2007). This compound has been employed to induce reversible inhibition of specific brain structures (Pelosi et al. 2007; Crestani et al. 2008; Resstel et al. 2008a; Tavares et al. 2009). It causes a reversible synaptic inactivation that spreads over an area of ~0.1–2 mm2. CoCl2 works by reducing Ca2+ pre-synaptic influx thus interfering with neurotransmitter release, causing a synaptic blockage without interfering with fibers of passage (Hagiwara and Byerly 1981; Lomber 1999). Thus, by causing temporary inactivation of local neurotransmission, the use of CoCl2 can minimize several problems associated with irreversible lesion techniques. Moreover, nonselective lesions of the LSA adjacent structures’ medial septal area (MSA) and bed nucleus of the stria terminals (BNST), which send projections to the hippocampus (Amaral and Kurz 1985; Everitt and Robbins 1997; Dong et al. 2001), produce profound deficits in spatial memory (Mizumori et al. 1990; Givens and Olton 1994; Everitt and Robbins 1997; Calandreau et al. 2007; Fitz et al. 2008). Therefore, the contradictory results observed after LSA irreversible lesions compared to the acute reversible synaptic blockage used in this study could be explained not only by the different functional impact of the experimental procedures, but also by a possible involvement of LSA adjacent structures.

Different from the present results, acute LSA inhibition by local injection of lidocaine before a conditioning session was reported to reduce freezing in mice re-exposed to an aversive conditioned context (Calandreau et al. 2007). There are, however, important differences between this latter study and ours. In addition to the distinct species used, lidocaine also affects fibers of passage by blocking voltage-gated Na+ channels (Lomber 1999). Moreover, in the previous study by Calandreau et al. (2007), a discrete stimulus (a tone) was included in the contextual fear conditioning session. Phillips and LeDoux (1994) showed that the dorsal hippocampus is engaged in the expression of contextual fear when a discrete cue is added to the context where the conditioning took place. In this situation the context is thought to predict the occurrence of the discrete cue (Phillips and LeDoux 1994).

The fimbria-fornix system is one of the main sources of afferent and efferent fibers of the hippocampal formation (Andersen et al. 1979). Despite some conflicting results, this system is proposed to play a key role in contextual fear response. Contextual fear deficits have been reported in rats with either fimbria-fornix or dorsal hippocampal lesions (Phillips and LeDoux 1995; Maren and Fanselow 1997). Electrophysiological studies have also suggested that the fimbria–LSA pathway is involved in this process. In mice, re-exposure to contextual conditioned stimuli has been associated with a decrease in synaptic transmission in the fimbria–LSA pathway. Moreover, tetanic fimbrial stimulation–induced long-term potentiation (LTP) in the lateral septum impaired the expression but not the acquisition of contextual fear conditioning (Vouimba et al. 1999). Low-frequency stimulation of this pathway, however, suppressed the impairing effect of tetanus on contextual conditioning and enhanced freezing to contextual stimuli. It was suggested that the level of hippocampal–septal neurotransmission and the magnitude of freezing would be related whenever the situation offered no possibility to predict and/or to avoid the aversive event (Vouimba et al. 1999).

The LSA role in organizing physiological, in addition to behavioral, components of defensive responses suggests its association with integrative centers controlling autonomic functions (Whitehead et al. 2000). In fact, the LSA has reciprocal connections with the central nuclei of the amygdala and receives projections from the hippocampus (Volz et al. 1990; Sheehan et al. 2004), the vMPFC (Vertes 2004), and BNST (Risold and Swanson 1997), brain areas with an established role in behavior and cardiovascular responses associated with contextual fear conditioning (Kim and Fanselow 1992; Phillips and LeDoux 1992; Resstel et al. 2006a, 2008a,b,c). Our results show that LSA inhibition before testing attenuated not only the behavioral but also the cardiovascular changes associated with exposure to the aversive context. This finding corroborates a large number of studies indicating that the LSA can modulate cardiovascular activity (Scopinho et al. 2006, 2007, 2008). Supporting this LSA role, it was described that its inactivation reduced the cardiovascular response observed in rats during acute restraint stress (Kubo et al. 2002) and the baroreflex activity in normotensive unanesthetized rats (Scopinho et al. 2007). Several neurotransmitter systems seem to be involved in these effects. Local LSA administration of noradrenergic and cholinergic receptor agonists increases blood pressure, whereas glutamate agonists cause vasodilation of hindlimbs. Both responses are observed during stress situations. Direct injection of GABA agonists into the LSA, on the other hand, reduces the cardiovascular responses associated with defensive behavior (Pérez-Polón and Corrêa et al. 1992; Kubo et al. 2002; Scopinho et al. 2006).

In conclusion, the present results indicate that the LSA could play an important role in the expression of fear conditioning, modulating both the behavioral and cardiovascular responses induced by re-exposure to the averesively conditioned context. They also suggest that this region is not essential for the acquisition and consolidation of contextual fear conditioning.

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