1 Enhancement of central norepinephrinergic neurotransmission contributes to

- 2 inhibition of seizure-induced respiratory arrest by targeting the beta-1 adrenergic
- 3 receptor (β 1-AR) locating in the cardiomyocytes in the DBA/1 mouse SUDEP model
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11 Abstract

12 Sudden unexpected death of epilepsy (SUDEP) is the key cause of of death in patients with epilepsy. Due to the complicated pathogenesis of SUDEP, however, the 13 exact mechanism of SUDEP remains elusive. Currently, although it is recognized that 14 the seizure-induced respiratory arrest (S-IRA) may be a main cause for SUDEP, other 15 16 factors resulting in SUDEP can not be excluded e.g arrhythmias. Our previous findings indicated that the incidence of seizure-induced respiratory arrest S-IRA and 17 SUDEP evoked by acoustic stimulation or pentetrazol (PTZ) injection was 18 significantly reduced by atomoxetine, a norepinephrine reuptake inhibitor (NRI), 19 20 suggesting that noradrenergic neurotransmission modulates S-IRA and SUDEP. Given that norepinephrine acts on the central and peripheral target to modulate respiratory 21 and circulation function by targeting adrenergic receptor α and beta (a-AR and β -AR) 22 and the arrhythmias can be contributed to SUDEP. Meanwhile, to further test whether 23 24 cardiac factors are implicated in S-IRA and SUDEP, we choose esmolol hydrochloride, a selective antagonist of beta-1 adrenergic receptor (β 1-AR) to test it in our models. 25 Our findings demonstrated that the lower incidence of S-IRA and SUDEP evoked by 26 acoustic stimulation or PTZ in DBA/1 mice by administration with atomoxetine was 27 significantly reversed by intraperitoneal (IP) of esmolol hydrochloride. Importantly, 28 29 the data of electrocardiogram (ECG) showed that the cardiac arrhythmia evoked by acoustic stimulation including the ventricular tachycardia, ventricular premature beat 30 1

and atrioventricular block and administration of atomoxetine significantly reduced theses arrhythmias and the incidence of S-IRA and SUDEP in our models. Thus, the dysfunction of respiratory and circulation may be implicated in the pathogenesis of S-IRA and SUDEP hand in hand and enhancing central norepinephrinergic neurotransmission contributes to inhibition of seizure-induced respiratory arrest by targeting β 1-AR locating in the cardiomyocytes. Our findings will show a new light on decoding the pathogenesis of SUDEP.

Keywords: sudden unexpected death in epilepsy (SUDEP); seizure-induced
respiratory arrest S-IRA); esmolol hydrochloride (Esmolol); Electrocardiogram
(ECG); locus coeruleus (LC); cardiac arrhythmia; pentetrazol (PTZ)

11 **1. INTRODUCTION**

12 It had been recognized that sudden unexpected death in epilepsy (SUDEP) is the key cause leading to the morality of epilepsy patients, especially those who had 13 undergone antiepileptic drug resistance ^[1-4]. Although some important advancements 14 concerning the pathogenesis of SUDEP had been obtained, to our certain knowledge, 15 16 decoding SUDEP seems to be a huge challenge in views of the complicated mechanism of SUDEP. Indeed, seizure-induced respiratory arrest (S-IRA) is an 17 important cause resulting in the death of epilepsy including our previous findings ^[2-6]. 18 However, more and more evidence hinted that the mechanisms of SUDEP are likely 19 20 to be heterogeneous ^[5]. Some data showed that the dysfunction in circulation may be a contributor to SUDEP. In fact, both of the dysfunction of respiratory and circulation 21 was in implicated in S-IRA and SUDEP. Previous findings showed that the cardiac 22 and respiratory function were simultaneously suppressed to induce the S-IRA and 23 24 SUDEP evoked by acoustic stimulation in DBA/1 mice, suggesting that the autonomic nerve including the sympathetic and parasympathetic nerve were involved 25 in the course of S-IRA and SUDEP. 26

Our previous data unveiled that the incidence of S-IRA and SUDEP evoked by acoustic or PTZ can be significantly reduced by IP with atomoxetine, a norepinephrine reuptake inhibitor (NRI) ^[6]. Considering that the central and peripheral norepinephrine transmission was associated closely with the activity of the

sympathetic nerve. It was well known that norepinephrine transmission modulates the 1 respiratory and circulation function by targeting the adrenergic $\alpha 1$, $\alpha 2$, $\beta 1$, and $\beta 2$ 2 receptors. The findings from our and other laboratories showed $\alpha 1$ and $\alpha 2$ receptors 3 were implicated in the S-IRA and SUDEP medicated by atomoxetine in the DBA/1 4 mice models ^[7-8]. However, whether both of β 1 and β 2 receptors, especially β 1 5 receptor located in the cardiomyocytes that play a key role in modulating the cardiac 6 function seems to essentially be explored based on our and other findings. Therefore, 7 8 we choose esmolol hydrochloride, a selective antagonist of beta-1 adrenergic receptor $(\beta 1-AR)$ to test whether the $\beta 1-AR$ was implicated in the atomoxetine mediated 9 S-IRA and SUDEP. Our findings showed that the lower incidence of S-IRA and 10 SUDEP evoked by acoustic stimulation medicated by atomoxetine was significantly 11 12 reversed by Esmolol. Thus, our data further demonstrated that the cardiac factors implicated in the reduction of the incidence of S-IRA and SUDEP by atomoxetine 13 can't be excluded and further contribute to explaining the possibility of heterogeneous 14 factors leading to SUDEP. 15

16 2. MATERIALS AND METHODS

17 2.1 Animals

All experimental procedures were the agreement with the National Institutes of Health 18 Guidelines for the Care and Use of Laboratory Animals and approved by the Animal 19 20 Advisory Committee of Zhejiang University. DBA/1 mice were housed and bred in the Animal Center of Zhejiang University School of Medicine and provided with 21 rodent food and water ad libitum. DBA/1 mice of either sex were used in this 22 experiment, as previous studies have shown, gender is not a variable that affects the 23 24 S-IRA of DBA/1 mice. Aiming at acoustic stimulation murine model, DBA/1 mice were "primed" starting from postnatal days 26-28 to establish the consistent 25 susceptibility to audiogenic seizures and S-IRA and the PTZ-evoked S-IRA model 26 was established as previously described ^[6-7]. 27

28 **2.2 Seizure induction and resuscitation**

S-IRA was established by acoustic stimulation as we previously described ^[6-7]. For
 the acoustic stimulation model, each mouse was placed in a cylindrical plexiglass
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chamber in a sound-isolated room, and audiogenic seizures (AGSZs) were evoked by 1 using an electric bell (96 dB SPL, Zhejiang People's Electronics, China). Acoustic 2 stimulation was given for a maximum duration of 60 s or until the onset of tonic 3 seizures and S-IRA in most mice in each group. Mice with S-IRA have resuscitated 4 within 5 s post the final respiratory gasp by a rodent respirator (YuYan Instruments, 5 Shanghai, China). S-IRA was also established in all non-primed DBA/1 mice by IP 6 administration of a single dose of PTZ (Cat #P6500; Sigma-Aldrich, St. Louis, MO) 7 at a dose of 75 mg/kg. 8

9 2.3 Effects of IP administration of Esmolol on the atomoxetine-mediated 10 suppression of S-IRA evoked by acoustic stimulation

The susceptibility of DBA/1 mice to S-IRA was confirmed 24 hours before 11 12 atomoxetine or vehicle administration. To IP administration, atomoxetine and esmolol were dissolved in saline. Different dosages of esmolol were applied to a vehicle 13 control group and treatment groups, in which DBA/1 mice would be acoustically 14 stimulated later. Atomoxetine (15 mg/kg, Ca # Y0001586; Sigma-Aldrich) or vehicle 15 16 was intraperitoneal (IP) injection in DBA/1 mice 120min prior to acoustic stimulation, 115min after infusion of atomoxetine (or vehicle), IP injection of esmolol (25mg/kg, 17 50mg/kg, H19991058; Qilu Pharmaceutical Co., Ltd.) or vehicle 5min before acoustic 18 stimulation (n=5-7/per group). The incidence of S-IRA, latency to AGSZs, duration of 19 20 wild running, clonic seizures, tonic-clonic seizures, and seizure scores were videotaped for offline analysis. DBA/1 mice from the different pre-treatment groups, 21 the monitoring of ECG was performed before acoustic stimulation and post-S-IRA. 22 To exclude the effects of esmolol on the incidence of the morality of DBA/1 mice, the 23 group, subjected to confirmation of S-IRA before 24 hours to experiment, was 24 pre-treated with the esmolol (50mg/kg, i.p) without administration of atomoxetine and 25 acoustic stimulation in the same manner to observe the incidence of the morality of 26 DBA/1 mice (n = 6-7/per group). 27

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2.4 Effects of IP administration of Esmolol on the atomoxetine-mediated suppression of S-IRA evoked by PTZ

The experiments were performed as follows: 1) For the control group, 3 saline (i.p) was administered 120 min prior to and 10 min post PTZ (75 mg/kg, i.p), respectively 4 (n=8); 2) For the Esmolol group, saline (i.p) was administered 120 min and prior to 5 and Esmolol (50 mg/kg, i.p.) was administered 10 min post PTZ (75 mg/kg, i.p), 6 respectively (n=7); 3) For the atomoxetine group, atomoxetine (15 mg/kg, i.p.) was 7 8 administered 120 min and prior to and saline (i.p) was administered 10 min post PTZ (75 mg/kg, i.p), respectively (n=5); 4) For the atomoxetine with Esmolol group, 9 atomoxetine (15 mg/kg, i.p.) was administered 120 min prior to and Esmolol (50 10 mg/kg, i.p) was administered 10 min post PTZ (75 mg/kg, i.p), respectively (n=5). 11 12 The S-IRA latency to AGSz, duration of wild running and clonic seizures, duration tonic-clonic seizures, and seizure scores were determined by offline analysis of video 13 recordings (n = 6-7/per group). 14

15 **2.6 Data analysis**

Statistical analyses were performed using SPSS 23(SPSS Software Inc., USA). The incidence of S-IRA was compared in different groups using Wilcoxon Signed Rank test, as these data are nonparametric. The data on seizure scores and the latency to AGSZs, the duration of wild running, clonic seizures, and tonic-clonic seizures were evaluated using the one-way ANOVA tests, which were presented as the mean \pm SEM. Statistical significance was inferred if p < 0.05.

22 **3. Results**

3.1 Administration of Esmolol significantly reversed the atomoxetine-mediated suppression of S-IRA evoked by acoustic stimulation

To investigate the effects of Esmolol on the incidence of S-IRA and SUDEP medicated by atomoxetine based on our previous findings. The following experiments were performed. The incidence of S-IRA evoked by acoustic stimulation in primed DBA/1 mice was obviously reduced by atomoxetine(15 mg/kg,i.p)compared with the vehicle group(n=6, n=7, p < 0.001). There was no significant difference between the vehicle group and the group pre-treated with esmolol (50 mg/kg,i.p) and with vehicle 5

(n=7,n=6, p > 0.05). However, the incidence of -S-IRA in the group pre-treated with 1 atomoxetine(15 mg/kg,i.p) and esmolol(50 mg/kg,i.p) was significantly increased than 2 the group pre-treated with atomoxetine(15 mg/kg,i.p) and the vehicle(n=6, n=7, p < 3 0.05). The incidence of -S-IRA in the group pre-treated with atomoxetine(15 4 mg/kg,i.p) and the vehicle was significantly decreased than in the group pre-treated 5 with the vehicle and with Esmolol(50 mg/kg,i.p) (n=6, n=6, p < 0.001). The 6 difference of latencies to AGSZs, durations of wild running, clonic seizures, 7 8 tonic-clonic seizures, and seizure scores in the treatment group showed administration 9 of esmolol significantly reversed the atomoxetine-mediated suppression of S-IRA evoked by acoustic stimulation without sensitivity to seizures. These data showed the 10 atomoxetine-mediated suppression of S-IRA evoked by acoustic stimulation can be 11 12 significantly reversed by esmolol, which means that administration of atomoxetine reduced the incidence of S-IRA by targeting the B1-AR locating in the 13 cardiomyocytes in our models. (FIG 1) 14

3.2 Administration of Esmolol specifically reversed the atomoxetine-mediated suppression of S-IRA by targeting the β1-AR locating in the cardiomyocytes

To exclude whether the dose of Esmolol (50 mg/kg,i.p) produces the morality of 17 DBA/1 mice to affect the specificity to reverse the reversed atomoxetine-mediated 18 suppression of S-IRA in our models. The following experiments were performed in 19 20 different groups. There was no significant difference in the morality of DBA/1 mice between the group pre-treated with the vehicle and the group pre-treated with Esmolol 21 (50 mg/kg,i.p) (P > 0.05), suggesting that the atomoxetine-mediated suppression of 22 S-IRA can be specifically reversed by esmolol by targeting the β 1-AR locating in the 23 24 cardiomyocytes. (FIG 2)

3.3 Obvious cardiac arrhythmia immediately occurred with tonic seizures and S-IRA evoked by acoustic stimulation

To observe the changing of ECG in the different treatment groups to analyze the role of atomoxetine and/or esmolol in cardiac electrophysiology. We observed that the ventricular tachycardia, ventricular premature beat, and atrioventricular block frequently occurred immediately following tonic seizures and S-IRA evoked by 6

acoustic stimulation in DBA/1 mice in a group pre-treated with the vehicle and with 1 2 the vehicle. However, Compared with the vehicle group, the arrhythmia was significantly decreased in the group pre-treated with atomoxetine (15 mg/kg,i.p) and 3 vehicle. We also observed the arrhythmia frequently occurred in the group pre-treated 4 with esmolol and the vehicle and the group pre-treated with atomoxetine (15 5 mg/kg,i.p) and esmolol, suggesting that Esmolol can specifically reverse the 6 atomoxetine-mediated suppression of S-IRA by targeting the β 1-AR locating in the 7 8 cardiomyocytes without resulting in extra arrhythmia. (FIG 3-4)

9 3.4 Administration of Esmolol significantly reversed the atomoxetine-mediated 10 suppression of S-IRA evoked by PTZ

To further investigate whether administration of Esmolol significantly reversed the 11 12 atomoxetine-mediated suppression of S-IRA depends on S-IRA and SUDEP models or not. The PTZ-induced S-IRA was accepted to test in the present study. Compared 13 with the vehicle group, the lower incidence of S-IRA by PTZ was reversed by 14 atomoxetine (p < 0.01). No obvious difference in the group of vehicle and in the group 15 of Esmolol (p>0.05), meaning that administration of Esmolol (50 mg/kg,i.p) 16 produced no effects on the incidence of mortality caused by PTZ itself. However, 17 Compared with the group treated with atomoxetine and vehicle, the incidence of 18 S-IRA and SUDEP significantly increased in the group treated with atomoxetine and 19 20 Esmolol (p < 0.01), suggesting that administration of Esmolol can specifically reverse the atomoxetine-mediated suppression of S-IRA evoked by PTZ. Thus, it can be 21 speculated from our findings that enhancing the norepinephrinergic neurotransmission 22 contributes to inhibition of S-IRA via targeting the $(\beta 1-AR)$ localized in the 23 24 cardiomyocytes in the DBA/1 mouse SUDEP model.

25 4. DISCUSSION

There are about 70 million people in the world who suffer from epilepsy, and 2.4 million have been diagnosed with epilepsy every year, nearly 90% occurred in developing countries^[1]. The latest epidemiological studies of the American Epilepsy Society show that the incidence of SUDEP in western developed countries is about 1.11 per 1000 people per year^[2], while the results of the multi-center epidemiological 7

survey in China show that the incidence of SUDEP in China is about 2.3 per 1000 1 people per year^[3], and the number of domestic epilepsy patients dying from SUDEP is 2 more than 20,000 per year. More importantly, SUDEP is the main factor that leads to 3 the death in young patients with epilepsy with an incidence two to three times that of 4 other diseases that can lead to sudden death in young adults, and children's SUDEP 5 incidence is also close to adults^[4]. Japan's latest epidemiological survey shows that 6 the incidence of SUDEP in pregnant women is significantly higher than the incidence 7 8 of non-pregnant women^[5]. However, due to the complexity of its pathogenesis and the difficulty to obtain the real-time monitoring data of SUDEP in patients with epilepsy, 9 the understanding of the SUDEP mechanism is still limited, and it is difficult to 10 develop a prevention strategy. Many previous studies have shown that the 11 12 pathophysiology of SUDEP is complex, including the disfunction of heart, autonomic nerves, respiratory, brain structural obstacles, and polygenesis. 13

In the current study, we firstly reported that the suppressant effects of S-IRA and 14 SUDEP in DBA/1 mice by atomoxetine can be significantly reversed by 15 16 administration of esmolol, a selective $\beta 1$ receptor blocker. Our previous findings showed that administration of atomoxetine inhibited S-IRA and SUDEP in DBA/1 17 mice independently of seizure induction methods, and its inhibition of S-IRA induced 18 by acoustic stimulation was dose-dependent^[6]. Given the current clinical reports that 19 20 atomoxetine can produce the effects of peripheral circulation. For example, after oral administration of atomoxetine, ADHD (Attention-Deficit/ Hyperactivity Disorder) 21 patients showed symptoms such as increased heart rate and other symptoms^[7], 22 considering that atomoxetine may play a role in SUDEP passing through the central -23 24 peripheral - cardiac sympathetic mechanisms by affecting $\beta 1$ receptors as dominate receptors in the heart. In the present study, we used the β 1 receptor blocker esmolol to 25 block the β 1 receptor which locates preferentially in the cardiomyocytes of the heart 26 to test whether the lower incidence of S-IRA by atomoxetine can be reversed by 27 Esmolol. It turned out that the suppressant effects of S-IRA by atomoxetine were 28 29 markedly reversed by esmolol, suggesting that administration of atomoxetine significantly reduced the incidence of S-IRA by enhancing the concentration of 30 8

norepinephrine of the cleft between the sympathetic synapse and cardiomyocytes of 1 the heart to combine with β 1 receptors of cardiomyocytes. Of course, atomoxetine 2 medicating the reduction of S-IRA by enhancing the concentration of norepinephrine 3 to target the adrenergic receptor $\alpha 1$ and $\alpha 2$ in the brain was the main pathway as well. 4 To be exact, atomoxetine medicates the reduction of S-IRA by interacting between $\alpha 1$, 5 $\alpha 2$, and $\beta 1$ receptors located in central and peripheral sympathetic nerves. In fact, 6 although S-IRA was the key cause leading to SUDEP, other factors are involved in the 7 8 course can't be excluded as well. A previous study showed that S-IRA evoked by acoustic stimulation in DBA/1mice was characterized by the simultaneous 9 suppression of respiration and circulation though the respiratory arrest preceded the 10 cardiac arrest. What's more, the recent study showed that the tonic phase apnea 11 12 evoked by acoustic stimulation is not sufficient for seizure-induced death in a W/+Emx1-Cre mouse. Therefore, the factors resulting in the S-IRA and SUDEP 13 including respiration and circulation may be heterogeneous. 14

The previous research has shown that the main risk factors of SUDEP are 15 16 generalized tonic-clonus and nocturnal seizures. The neuronal network of generalized tonic-clonic seizures may lead to hypoventilation, apnea, and cardiovascular failure 17 by inhibiting brainstem respiration or autonomic nerve control centers, leading to 18 death ^[10, 11]. Since the most sudden death cases occur at night or in sleep, with no 19 20 witnesses, leaving many questions unanswered. In recent years, more and more researches have shown that apnea and heart failure are important potential 21 mechanisms in SUDEP. Studies in animal models and SUDEP patients have shown 22 that seizure-induced respiratory arrest (S-IRA) is one of the main induced factors that 23 lead to death in many cases^[12-14]. However, a respiratory arrest is not the only lethal 24 cause, tonic-clonic seizures caused sudden, simultaneous respiratory and cardiac 25 depression in mice, and the time of recovery of respiration and heart rate during 26 resuscitation is also closely matched^[15]. Therefore, the heart also plays an important 27 role in the mechanism of SUDEP. 28

Seizure-associated arrhythmias are common and have been considered as the
 potential pathogenesis of SUDEP, such as tachycardia, bradycardia, and cardiac arrest
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after the seizures, resulting in cardiac repolarization, changes in electrolyte levels, 1 blood pH, and catecholamine release in patients with epilepsy, which latter three 2 factors may promote arrhythmias and lead to death by affecting cardiac excitability^[16]. 3 Tachycardia and several factors that increase susceptibility to tachyarrhythmias, 4 known as pathological cardiac repolarization, are considered as risk factors for sudden 5 cardiac death (SCD) in healthy individuals and patients with epilepsy which, in this 6 particular setting, are important factors in triggering SUDEP events^[17, 18]. However, 7 8 the exact mechanism of SUDEP and cardiac arrest induced by seizures remains unclear. Seizures may interfere with the normal functioning of the heart by causing 9 autonomic disturbances ^[19], leading to fatal arrhythmias. Studies have found that 10 cardiac electrical instability and autonomic dystonia are caused by cumulative cardiac 11 damage during repeated attacks^[20]. Studies also have shown that catecholamine 12 (norepinephrine and adrenaline) released from seizures is related to a wide range of 13 calcium (Ca2+)-mediated physiological changes, which can lead to damage to the 14 heart structure. Our previous studies found that repeated S-IRA in DBA / 1 mice will 15 16 result in necrotic damage in the ventricle which is caused by the local Ca2 + homeostasis disturbances, and the incidence and severity of injury depend on the total 17 number of S-IRA^[21]. Relevant clinical data have found that paroxysmal arrhythmias 18 (including arrest, atrioventricular block, and rare atrial and ventricular fibrillation) 19 usually occur after convulsive episodes and are often associated with (near)SUDEP^[22]. 20 Studies have shown that chronic epilepsy can lead to hypoxemia and the increase of 21 catecholamine, causing damage of the heart and vascular, resulting in cardiac 22 electrical, and mechanical dysfunction^[23,24]. Some fatal incentives may be the acidosis 23 caused by persistent hypoxemia and hypercapnia, leading to bradycardia or cardiac 24 arrest, while in others it may be the seizures due to malignant arrhythmias^[25]. This 25 suggests that SUDEP is inextricably linked to arrhythmia. 26

In the present study, the ECG of DBA/ 1 mice with administration of esmolol (50 mg/kg, i.p) that can significantly reverse the lower incidence of S-IRA evoked by acoustic stimulation by administration of atomoxetine showed the mixture of sinus bradycardia, atrioventricular block, ventricular premature beat, and ventricular

tachycardia in order. In the meantime, the ECG of DBA/1 mice suffered from the 1 S-IRA evoked by acoustic stimulation characterized by the mixture of sinus 2 bradycardia, atrioventricular block, ventricular premature beat, and ventricular 3 tachycardia as well in our model. However, no obvious arrhythmia appeared in the 4 DBA/1 mice without suffering from S-IRA other than the sinus bradycardia and no 5 apparent mortality occurred in the group of DBA/1 mice with pre-treatment with the 6 dose of esmolol 50 mg/kg (i.p). Furthermore, the arrhythmias can be significantly 7 8 reduced by atomoxetine in our models. Thus, the lower incidence of S-IRA by acoustic stimulation or PTZ can be significantly as well as especially reversed by 9 esmolol by blocking the B1-AR localized in cardiomyocytes to combine the 10 norepinephrine released by cardiac sympathetic nerve synaptic terminal in our models. 11 12 Additionally, it is possible that the locus coeruleus (LC), as the largest nuclei to synthesize and release norepinephrine in the brain, may be implicated in the course of 13 administration of esmolol to reverse the lower incidence of S-IRA and SUDEP and 14 plays a key role in modulating S-IRA and SUDEP by regulating the respiratory and 15 16 circulation function in our models. In future experiments, we will further explore it.

17 5. Conclusions

The finding models 18 current in our suggested that enhancing central norepinephrinergic neurotransmission contributes to inhibition of seizure-induced 19 20 respiratory arrest by targeting β 1-AR locating in the cardiomyocytes and the pathway of norepinephrinergic neurotransmission- β 1-AR between brain and heart may be a 21 potential and specific target to prevent SUDEP. 22

23 6. Funding

The work was supported by the National Natural Science Foundation of China (Grant.NO: 81771403 and 81974205); by the Natural Science Foundation of Zhejiang Province (LZ20H090001); by the Program of New Century 131 outstanding young talent plan top-level of Hang Zhou to HHZ

28 7. DISCLOSURE

29 All authors declare no competing interests. We had confirmed that we have read the

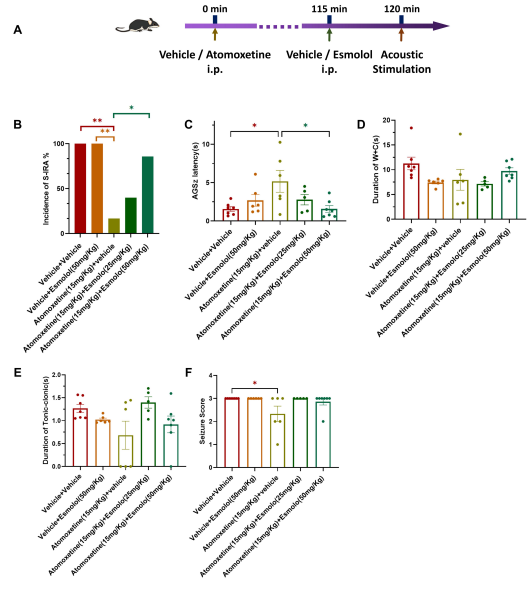
30 Journal's position on issues involved in ethical publication and affirm that this study 11

1 was in accordance with those guidelines.

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3 8. Figure legends





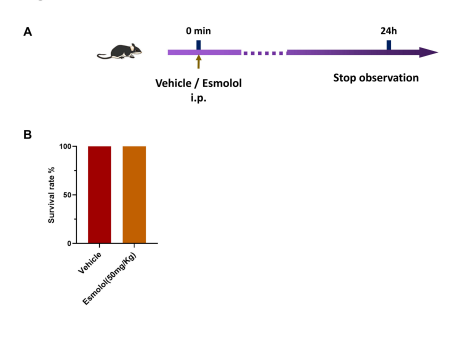
- 4 5
- Figure 1 Administration of Esmolol significantly reversed the atomoxetine-mediated
 suppression of S-IRA evoked by acoustic stimulation

A. The protocol to explore the influence of administration of esmolol on
atomoxetine-mediated reduction of S-IRA evoked by acoustic stimulation. B.
Compared to the group treated with vehicle (n = 7) or vehicle and esmolol (n = 6),
S-IRA evoked by acoustic stimulation was markedly lower in groups treated with i.p. 12

atomoxetine (n = 6, **p < 0.01) at 15mg/Kg. However, the protective effect of 1 atomoxetine in primed DBA/1 mice was significantly reversed by esmolol doses of 2 50 mg/kg (n = 7, *p < 0.05). C. Furthermore, compared with the group treated with 3 atomoxetine and vehicle (n = 6), the latencies to AGSZs in the control group (n = 7)4 or the group treated with atomoxetine and esmolol (n = 7) was significantly increased 5 (p < 0.05). **D-E.** There were no intergroup differences in durations of wild running 6 plus clonic seizures (W+C), tonic-clonic seizures (p > 0.05). F. Compared to the 7 8 control group (n = 7), the seizure score was lower in the group treated with 9 atomoxetine and vehicle (n = 6, *p < 0.05).







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Figure 2 The dose of Esmolol (50 mg/kg,i.p.) does not produce the incidence of
morality of DBA/1 mice.

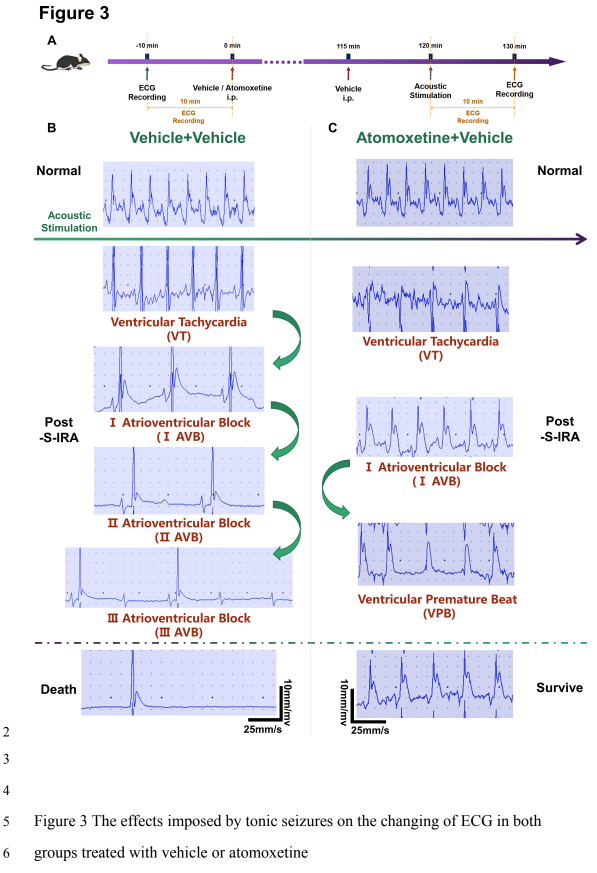
A. The protocol to explore the dose of esmolol (50 mg/kg,i.p.) whether produce the morality of DBA/1 mice. **B.** There was no significant difference in the morality of DBA/1 mice between the group pre-treated with the vehicle and the group pre-treated with esmolol(50 mg/kg, i.p, n=7) (p > 0.05)

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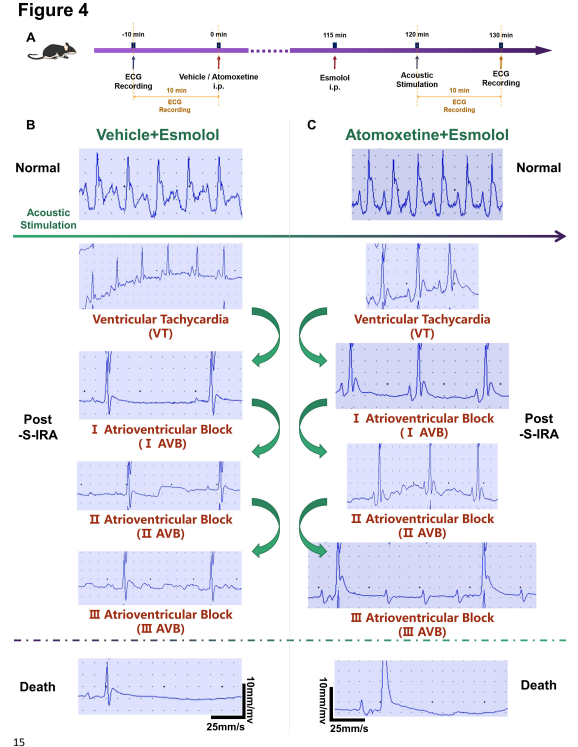
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7 A. The protocol to explore the changes of electrocardiogram on 14

atomoxetine-mediated reduction of S-IRA evoked by acoustic stimulation. **B.** The ventricular tachycardia and atrioventricular block occurred immediately following tonic seizures and S-IRA evoked by acoustic stimulation in DBA/1 mice in the control group. **C.** What's more, the group pre-treated with the atomoxetine (15mg/Kg) was observed the ventricular tachycardia, atrioventricular block, and ventricular premature beat after acoustic stimulation, and the DBA/1 mice were then recovered.



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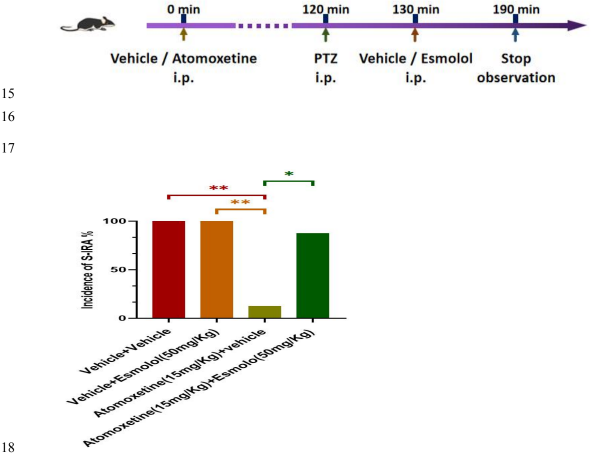
2 Figure 4 The effects imposed by tonic seizures on the changing of ECG in both

3 groups treated with atomoxetine or esmolol

A. The protocol to explore the changes of electrocardiogram on the administration of 4 esmolol reversed the atomoxetine-mediated reduction of S-IRA evoked by acoustic 5 stimulation. B. The ventricular tachycardia and atrioventricular block occurred 6 7 immediately following tonic seizures and S-IRA evoked by acoustic stimulation in 8 DBA/1 mice in the group treated with vehicle and Esmolol (50mg/Kg). C. The group 9 pre-treated with the atomoxetine(15mg/Kg) and treated with esmolol (50mg/Kg) was observed the ventricular tachycardia and atrioventricular block after acoustic 10 stimulation. 11

Figure 5 Administration of Esmolo significantly reversed the
atomoxetine-mediated suppression of S-IRA evoked by PTZ

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1 Compared with the group treated with vehicle (n = 5) or vehicle and esmolol (n = 5). 2 S-IRA evoked by acoustic stimulation was markedly lower in groups treated with i.p. atomoxetine (n = 7, **p < 0.01). However, Compared with the group treated with i.p. 3 atomoxetine and vehicle, the incidence of S-IRA significantly increased in the group 4 treated with i.p. atomoxetine and esmolol (n = 7, **p < 0.05) 5 9. References 6 SINGH A, TREVICK S. The Epidemiology of Global Epilepsy [J]. Neurol 7 [1] 8 Clin, 2016, 34(4): 837-47. 9 [2] LHATOO S, NOEBELS J, WHITTEMORE V, et al. Sudden unexpected death in epilepsy: Identifying risk and preventing mortality [J]. Epilepsia, 2015, 10 56(11): 1700-6. 11 12 [3] GE Y, DING D, ZHANG Q, et al. Incidence of sudden unexpected death in epilepsy in community-based cohort in China [J]. Epilepsy Behav, 2017, 76: 13 76-83. 14 KELLER A E, WHITNEY R, LI S A, et al. Incidence of sudden unexpected [4] 15 16 death in epilepsy in children is similar to adults [J]. Neurology, 2018, 91(2): e107-e11. 17 TANAKA H, KATSURAGI S, HASEGAWA J, et al. Maternal Death Related [5] 18 to Sudden Unexpected Death in Epilepsy: A Nationwide Survey in Japan [J]. 19 20 Brain Sci, 2021, 11(8). [6] ZHANG H, ZHAO H, FENG H J. Atomoxetine, a norepinephrine reuptake 21 inhibitor, reduces seizure-induced respiratory arrest [J]. Epilepsy Behav, 2017, 22 73: 6-9. 23 Yue Shen, Hai Xiang Ma, Han Lu 3, Hai Ting Zhao, Jian Liang Sun, Yuan 24 [7] Cheng, Hong Hai Zhang. Central deficiency of norepinephrine synthesis and 25 norepinephrinergic neurotransmission contributes 26 to seizure-induced arrest.Biomed Pharmacother. 2021:111024. 27 respiratory doi: 10.1016/j.biopha.2020.111024. 28 29 [8] Rui Zhang, Zheren Tan, Jianguo Niu Hua-Jun Feng. Adrenergic α2 receptors are implicated in seizure-induced respiratory arrest in DBA/1 mice. Life Sci. 30

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19