Efficacy of Low Dose Tenecteplase in Restoring Vessel Patency in a Clinically Relevant Large Animal Experimental Model

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Abstract

Background: We tested whether tenecteplase, a recombinant fibrin-specific plasminogen activator can induce efficient thrombolysis in lower doses than the hitherto indicated, compared to full-dose therapy for vessel thrombosis.

Methods: Farmed pigs (30-35 kg, n=23) were imposed in a clinically relevant model of common carotid-arteries (CA) and the left-anterior-descending (LAD) artery below the first diagonal occlusion. The vessels were initially externally injured followed by an appropriate for the vessel dose of thrombin. Occlusion was achieved in 80% for the LAD and 57% for the CA. Post occlusion the animals were allocated to one of the following groups: Group A) half-dose tenecteplase (2000IU); Group B) Full dose tenecteplase (5000IU); Group C) saline only. All animals received heparin and antiplatelet agents. The animals were followed-up for 120 minutes.

Results: Sixty minutes after thrombolysis recanalization occurred in 58% of the occluded arteries in Group A, in 85% in Group B and in 20% in Group C (P=0.028). Both tenecteplase regimens were effective for treating CA thrombosis (71% Group A vs 75% in Group B vs 33% in Group C, P=0.286), but low dose was significantly less effective in reestablishing flow in LADs (25% in Group A vs 100% in Group B vs 0% in Group C, P=0.015). Vessel patency was sustained for at least 1hr. In Group A, flow velocity in LAD reached 30% of the baseline value and 45% in carotids, while in Group B flow was 180% and 55% respectively. No differences in hemodynamic tested parameters and in post reperfusion arrhythmias were observed between the groups.

Conclusions: We developed a clinically relevant model useful for testing the efficacy of different thrombolytic regimens. Our data demonstrate that low dose tenecteplase can be successfully used for treating thrombosis of larger diameter vessels, such as the carotids, but appears inadequate for treating LAD thrombotic occlusions.

Keywords: Thrombolytics; Vessel thrombosis; Preclinical models

Introduction

Myocardial infarction is the leading cause of morbidity and mortality in western countries [1]. The most common cause is rupture of an atherosclerotic plaque with formation of a thrombus leading to occlusion of a coronary artery [2,3]. Primary percutaneous coronary intervention (PCI) is routinely performed in the current practice. However, even with advanced commute technological systems catheterization is not always performed timely and valuable time for myocardial salvage is spared.

Thrombolytic reagents emerged as a potential non-operative tool for early in situ recanalization; initial studies reported ante grade perfusion of the infarct related artery in 80% of the cases within 90 min, resulting in 50% reduction in mortality [4,5]. Additional advantages include wide availability, no need for institutional resources or experienced operator and the ability to deliver them in the emergency department or even before arriving to the hospital [6]. However, early enthusiasm was tempered by suboptimal reperfusion rates, recurrent ischemia and re-infarction and the most important, intracranial or systemic bleeding complications [7].

Among all the fibrinolytic drugs available, tenecteplase is of particular interest since it has the highest fibrin specificity and a long half-life (20-25min), allowing it to be given as a single-bolus dose. The first clinical trials, ASSENT2 [8] and ASSENT3 [9] proved tenecteplase equally safe and efficient to alteplase, while the latest clinical trial highlighted its role in comparison to PCI [10]. Here, we report our experience first with a highly efficient protocol of generating a clinically relevant model of vessel thrombosis and second that low dose of tenecteplase can be efficient in improving flow in large vessels but not in the LAD.

Methods

Animals

Experiments were performed on healthy domestic pigs of either sex weighing 30-35kg as depicted in Figure 1. A total of 23 animals were used for the current study. N=5 were used for the protocol optimisation, n=3 were excluded due to arrhythmias and n=5 were used in each experimental group. Abbreviation: TNKase= tenecteplase. The animals were premedicated by intramuscular administration of ketamine (20 mg/kg) and diazepam (1 mg/kg). Anesthesia was then induced and maintained by a continuous infusion of sufentanil and pentobarbital into an ear vein. The animal was intubated and ventilated with room air. Body temperature was maintained using a heating pad. Muscle relaxation was achieved by pancuronium bromide infusion (0.2 mg/kg).

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**Experimental procedure**

The right and left cervical area were incised to expose both common carotids (CCA) and the connective tissue around the arteries was dissected carefully. A ring-shaped transit time Doppler flow probe, suitable for the carotid artery diameter (5-8 mm), was placed proximally around the arteries, and mean blood flow was recorded continuously.

A segment of both common carotids 1 cm long was isolated distal to the flow probe with two Silastic vessel snares. Intimal injury was then created between the snares by four to five external compressions (one every 8-10 min) using hemostat forceps for about 30 minutes. The snares were then tied to occlude the CCA resulting in complete cessation of flow. Thrombin (190 UI) was mixed with blood and injected into the isolated segment. Thirty minutes later, the proximal Silastic snare was removed, immediately followed by the distal snare. Absence of carotid blood flow for at least 10 min after snare removal was considered as complete vessel thrombosis. A scheme of the experimental protocol and of the thrombosis model is depicted in Figures 2A and 2B.

The same procedure was applied to the left anterior descending artery just below the first diagonal (2-4 mm). Here a segment 0.5 cm long was isolated with two Silastic vessel snares carefully, after the first diagonal branch. Intimal injury was then created between the snares by three to four external compressions (one every 10-15 min) with a hemostat forceps for about 30 minutes. During this period severe cyanosis was developed in the subtended myocardial area followed by three to four external compressions (one every 10-15 min) with a hemostat forceps for about 30 minutes. The snares were then tied to occlude the CCA resulting in complete cessation of flow. Thrombin (190 UI) was mixed with blood and injected into the isolated segment. Thirty minutes later, the proximal Silastic snare was removed, immediately followed by the distal snare. Absence of coronary blood flow for at least 10 min after snare removal was considered as complete vessel thrombosis (Figure 2).

**Hemodynamic and electrocardiographic measurements:** Continuous electrocardiogram monitoring (ECG; lead II) and a flow meter were used for the evaluation of ST segment changes and recanalization respectively. Frequent Ventricular Premature Beat (VPB) was defined as more than 5 VPBs per minute. Accelerated Idioventricular Rhythm (AVIR) was defined as a ventricular rhythm with a rate of 60 to 125 beats/minute and frequent episodes of slow ventricular tachycardia. Nonsustained ventricular tachycardia (VT) was defined as 3 or more consecutive ventricular ectopic beats at a rate >100 beats/minute and lasting <30 seconds. Sustained VT was defined as the last longer than 30 seconds or cause hemodynamic compromise that requires intervention. Ventricular fibrillation was recognized by the presence of irregular nodulations of varying contour and amplitude. A Millar catheter was advanced through an incision at the LV apex in the LV chamber for continuous pressure monitoring. Complete or not complete resolution was defined with respect to initial ECG changes. Re-thrombosis was defined as a new ST-elevation compromised flow compared to the initial evaluation. After surgical preparation the animals were allowed 10 min for stabilization and then control measurements of ECG, flow and mean arterial pressure (MAP) were taken. Repeat measurements were taken again before occlusion; 10 min post successful thrombosis and before therapy administration; immediately after reperfusion and then every 30 min until the end of the study. Mean arterial pressure-rate product (PRP) was calculated as mmHg/beats/min \( \times 10^3 \).

**Antithrombotic strategy during the procedure:** All animals received antiplatelet and anticoagulation therapy according to the current recommendations. Oral aspirin 300 mg (purchased by Bayern) and oral clopidogrel 600 mg (purchased by Sanofi) at loading doses. Although the objective of the current study was to compare low and high dose of tenecteplase per se, in order to maximize the potential of recanalization high doses of antiplatelets were administered. Anticoagulation therapy including 60UI of unfractionated aspirin (purchased by Braun) was given intravenously. Unfractionated heparin instead of enoxaparin was used due to supply reasons.

**Thrombolytic therapy: dose of administration:** Post successful thrombosis of the vessels animals were allocated to low-dose (Group A) of intravenous tenecteplase (TNK-ase; purchased by Boehringer Ingelheim) slowly administered over 4-5min through jugular vein catheter, or high dose (Group B) or saline. The longer than usual time of infusion based on the half-life of tenecteplase should not interfere with the result.

**Biochemical studies**

In all groups, before euthanasia, blood samples were collected in glass citric tubes (BD Vacutainer) and processed according to manufacturer’s instructions. Coagulation markers such as international normalized ratio (INR), activated partial thromboplastin time (aPTT) and prothrombin time (PT) were evaluated.

**Statistical analysis**

The results are expressed as mean ± SEM, and the data were processed by factorial one-way ANOVA. Individual groups were compared using Dunnett’s t test. Differences with \( P<0.05 \) were considered statistically significant.

**Results**

A total of 23 animals were used in the current study. 5 animals were initially included for the thrombus formation optimization protocol. Another three animals developed ventricular fibrillation during thrombus formation and died before treatment administration. A total of 15 animals were included in the three threatened groups.

We decided to use an optimized protocol with thrombin infusion to interrupt vessel flow rather than mechanical interventions with percutaneous catheters so as to better simulate a clinical acute MI scenario. The latter entails a ruptured plaque, adhesion of platelets and thrombi formation. Endothelial disruption by external compressions and infusion of blood with thrombin mimicked this pathology. Results show that successful thrombosis was achieved in 80% of the carotids.
and in 57% of the LAD vessels used.

**Recanalization**

Ten minutes after the induction of occlusive thrombi, we tested whether the low or the high dose of TNK-ase are able to restore flow in the thrombosed vessels. To this end, 2000 IU TNK-ase (Group A, n=5) or 5000IU TNK-ase (Group B, n=5) or saline only (Group C, n=5), were injected via the internal jugular vein. All animals received 60IU/kg heparin and antiplatelet therapy. No adverse events were observed during the infusion in any of the animals studied. At 60 minutes post lytic drug administration, recanalization occurred in 58 ± 8% of the vessels in Group A, 85 ± 15% in Group B and 20 ± 5% in Group C (Figure 2C; p=0.028). Both tenecteplase regimens were effective for treating carotid artery thrombosis (Figure 2D; 71 ± 5% Group A vs 75 ± 4.5% in Group C, P=0.286), but low dose was significantly less effective in reestablishing flow in proximally occluded LADs (25 ± 6% in Group A vs 100% in Group B vs 0% in Group C, Figure 2E; p=0.015). All vessels with restored flow at 60 minutes remained patent at least for 1hr. In Group A, at 60 minutes after tenecteplase administration, flow velocity in LAD reached 30 ± 22% of the baseline value and 45 ± 12% in carotids, while in Group B 160 ± 42% and 55 ± 18% respectively (Figure 3A, B; p=0.028).

**Post-reperfusion arrhythmias**

ECG monitoring was performed during the whole experimental procedures for arrhythmias detection. No significant differences were observed in the arrhythmias observed during the vessel thrombosis. Previous studies showed that during the first 30 seconds of reperfusion there is a marked inhomogeneity in the action potentials within the ischemic area and at the border that contributes to the occurrence of fibrillation [11]. In line with these data we observed episodes of fibrillation in total 2 out of the 5 animals that required resuscitation within 30 min post reperfusion in Group A and in 4 animals in Group B. Episodes of AIVR were observed in Group C (2 out of 5 animals), but not in Group A or B. No statistically significant difference between

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**Figure 2:**
- **A.** Experimental protocol. **B.** Scheme of the protocol used to induce thrombosis. Following external damage to the endothelium, thrombin was injected and total occlusion was validated by a probe distal to the occlusion. **C.** Approximately 58 ± 8% of the vessels in Group A and 85 ± 15% in Group B were successfully recanalised compared to 20 ± 5% only in Group C. **D.** No difference between low and high dose Groups was observed as far as carotid patency is concerned. **E.** Significantly higher number of LADs were recanalised in the high dose of TNKase Group compared to the low dose or the Group that received only heparin. Abbreviation: TNKase= tenecteplase.

**Figure 3:**
- **A.** Percent of flow restoration in Carotids post high or low dose of TNKase. **B.** Percent of flow restoration in the LAD post high or low dose of TNKase. Abbreviation: TNKase=tenecteplase.
the groups in terms of sustained ventricular tachycardia, nonsustained ventricular tachycardia, frequent ventricular premature beats, and atrioventricular block was found. The association of ST segment regression and consequently higher rates of reperfusion arrhythmias was also evaluated. The animals in Group B revealed greater ST segment regression and consequently higher rates of reperfusion arrhythmias. The latter may be related to maximal heterogeneity when irreversibly and reversibly injured cells are juxtaposed.

**Hemodynamics**

Table 1 summarizes these data. Continuous LV pressure monitoring was performed with a Millar pressure catheter installed in LV apex. There was no statistically significant difference between LV systolic and diastolic pressure between all groups before treatment administration. Systolic blood pressure was significantly compromised in Group C compared to the other 2 groups. Failure to recanalize the LAD may explain this finding.

**Biochemical indexes**

No differences were observed in the INR values at the time point of evaluation (<1.2 in all animals evaluated) or aPTT or PT. Perhaps the short time post infusion of tenecteplase and heparin that these indexes were evaluated did not interfere with the systemic levels. Nevertheless, our findings are in line with previous reports were concomitant administration of thrombolytics and heparin did not influence acutely the hemostatic pathway [9,12,13].

**Discussion**

Restoration of epicardial coronary flow post-acute MI improves clinical outcome [14]. Recanalization can be achieved by means of thrombolysis or primary angioplasty. Although angioplasty is preferred by European and American guidelines accumulated studies question the superiority of angioplasty over thrombolysis [15,16]. The safety and effects and exerts higher thrombolytic potency even when given more than 4 hours after the onset of symptoms [8,20,22]. More than 25000 patients have been so far recruited in clinical trials that used tenecteplase as a potential therapeutic regimen making the use of this drug evidence based. In line with previous data we did not observe any changes in the biochemical indexes evaluated in the current study post either low or high dose of tenecteplase without changing the dose of heparin used in parallel [23]. However, low dose of tenecteplase restored blood flow in 25% of the LADs compared to 100% restored post full dose. Similar efficacy was observed between the two doses in regards to common carotid blood flow restoration, suggesting that low dose can be used safely for restoration of flow in large arterial vessels but not in the smaller in diameter ones.

The most common cause of MI is rupture of an atherosclerotic plaque with formation of a thrombus leading to occlusion of a coronary artery [2,3]. Serial *in vitro* experiments have examined the mechanism of atherosclerosis and have paved the way for new therapeutic drug development [17]. In addition, *in vivo* experiments in small genetically modified animals have also elucidated the pathology of the coronary syndrome [18]. However, the most suitable model should simulate the human disease. In the current study, we used a swine model to create thrombosis and mimic a clinical scenario of myocardial infarction. We first caused external damage to the common carotids and to the LAD so as to activate the aggregation of platelets and we consequently infused thrombin so as to create thrombosis. Platelet function is similar between swine and humans and this made them suitable for the current protocol [19]. In addition, coronary anatomy in swine is similar to the human except for left azygos vein which drains to the left coronary sinus. These data support the idea of a preclinical model of thrombosis in swine that would enable testing new therapeutic drugs for atherosclerosis. Indeed, following the protocol described in the current study we successfully occluded 80% of the LADs and 57% of the common carotids and then we tested tenecteplase efficacy to restore blood flow.

Different thrombolytic drugs have been used in preclinical and clinical studies for acute myocardial infarction with salutary effects reported. Among others tenecteplase is more promising given its high fibrin specificity that enables delivery by single bolus intravenous injection [20]. More specifically, tenecteplase is a genetically engineered plasminogen activator with reduced risk for causing major bleeds due to restricted lytic activity to the plasmin on the fibrin surface that avoids the breakdown of fibrinogen, factor V, factor VIII and a2-antiplasmin [21]. Compared to other thrombolytic regimens, tenecteplase is not affected by other drugs such as nitrates, it has more robust anti-platelet effects and exerts higher thrombolytic potency even when given for more than 4 hours after the onset of symptoms [8,20,22].

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**Table 1**: Hemodynamics during thrombosis formation experiments.

vessel patency in contrast to our findings. A possible explanation may be the smaller diameter of the LAD in our experimental model compared to the human atherosclerotic ones. More specifically, the diameter of the LAD used here, was approximately 2-3 mm while in humans it reaches 4-5 mm; the latter is closer to the swine carotid arteries that were recanalized with both low and high doses (~5 mm). In addition, although we used an experimental model of thrombus formation in order to simulate clinical atherosclerotic plaques, the external damage of the LAD can also result in vessel spasm and vessel internal diameter stenosis that may further aggravate the occlusion requiring higher doses of tenecteplase.

In conclusion, in the current study we successfully created a preclinical model of coronary thrombosis where different regimens can be routinely evaluated. Second, we used this model to test the efficacy of low and high dose of tenecteplase to restore blood flow in the coronaries and the common carotids. We found significant increase in the number of LADs treated with low dose while in common carotids both regimen doses were equally efficient. Following pioneering clinical trials [24,25] where thrombolysis by means of tenecteplase was very competitive with primary angioplasty when delivered very rapidly, possibly in the pre-hospital phase our data support that tenecteplase may offer a very simple and effective treatment, particularly if subsequent primary angioplasty is offered to the patients with high clinical risk or with recurrent ischemia.

References


