Enhanced clot lysis by continuous-wave ultrasound exposure an *in vitro* study using streptokinase and reteplase

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Abstract

Background
Ultrasound (US) has been used to enhance thrombolytic therapy in the treatment of stroke. Considerable attenuation of US intensity is however noted if US is applied over the temporal bone, the aim of this study was therefore to explore possible changes in the effect of thrombolytic drugs during low-intensity, high-frequency continuous-wave ultrasound (CW-US) exposure.

Methods
Clots were made from fresh venous blood drawn from healthy volunteers. Each clot was made from 1.4 ml blood and left to coagulate for 1 hour in a plastic test-tube. The thrombolytic drugs used were, 3600 IU streptokinase (SK) or 0.25 U reteplase (r-PA), which were mixed in 160 ml 0.9 % NaCl solution. Continuous-wave US exposure was applied at a frequency of 1 MHz and intensities ranging from 0.0125 to 1.2 W/cm². For each thrombolytic drug and US intensity, six clots were immersed in thrombolytic solution and exposed to US, while six control clots were exposed to thrombolytic solution alone. To evaluate the effect on clot lysis, the haemoglobin (Hb) released from each clot was measured every 20 min for 1 hour (20, 40 and 60 min). The Hb content (mg) released was estimated by spectrophotometry at 540 nm. The difference in effect on clot lysis was expressed as the difference in the amount of Hb released between pairs of US-exposed clots and control clots. Statistical analysis was performed using Wilcoxon’s signed rank test.

Results
Ultrasound significantly decreased the effects of SK at intensities of 0.9 and 1.2 W/cm² at all times (P < 0.05). Ultrasound significantly increased the effects of r-PA on clot lysis following 20 min CW-US exposure at 0.9 W/cm² and at 1.2 W/cm².
following 40 min exposure at 0.3, 0.6, 0.9 and at 1.2 W/cm$^2$, and following 60 min of CW-US exposure at 0.05 0.3, 0.6, 0.9 and at 1.2 W/cm$^2$ (all P < 0.05).

**Conclusions**
Increasing intensities of CW-US exposure resulted in increased lysis of r-PA-treated blood clots, but decreased clot lysis of SK-treated clots.

**Background**
Ultrasound (US) has been used to enhance thrombolytic therapy, for example, in the treatment of stroke. In this setting, US is usually applied over the temporal bone, exposing the obstructed vessel to US concomitantly with treatment with thrombolytic drugs [1-3]. The enhancement of various thrombolytic drugs has been demonstrated during *in vitro* clot lysis at frequencies, ranging from 20 kHz to 4.5 MHz [4-7]. Positive effects on clinical outcome have been reported when high frequency US has been used *in vivo* [1, 8]. However, in the CLOTBUST trial, the effects were found not to be statistically significant [1]. This is in contrast to the results from *in vitro* studies of US-enhanced thrombolysis, where considerable enhancement effects of the clot lysis have been shown as a result of exposure to US [7, 9, 10]. This discrepancy might be explained by the attenuation of US intensity passing through the temporal bone structure during high frequency US exposure. There have been reports of decreases in the output intensity between 86.8 % and 99.2 % when US is applied over the temporal bone [11, 12]. Low frequency US on the other hand, has greater penetration through bone tissue compared to high frequency US, resulting in higher US intensities reaching the obstructed vessel [13, 14]. However, low frequency US has been shown to induce a higher rate of bleeding complications during US-enhanced thrombolysis *in vivo* [15].
Other factors of US than intensity and frequency also seem to affect the results during US-enhanced thrombolysis. We have previously only found, during pulsed-wave US exposure that SK induced clot lysis, enhanced effects at low intensity (0.5 W/cm²) [16, 17]. During pulsed-wave US exposure of r-PA induced clot lysis, enhancement effects occur both at high and low intensity (i.e. ≤ 0.25 W/cm² or > 2.0 W/cm²) [18]. The enhancement effects might thus depend on duty cycle, i.e. the number of pulses sent [9, 19, 20]. Meunier et al reported increasing effects on tissue type plasminogen activators mediated clot lysis depending on increasing duty cycle [20]. However, Holland et al failed to verify the same duty cycle dependency [9]. Others have shown higher grades of enhancement using CW-US exposure than when pulsed-wave US exposure was used [5, 21].

The aim of this study was to investigate the changes in the effect of clot lysis of r-PA and SK during low-intensity, high-frequency CW-US exposure, intensities within the area following attenuation from the skull bone.

Methods
The methods employed for clot formation and clot lysis evaluation, and the ultrasonic properties of the model have been described in detail previously [17, 18, 22]. Only a brief description will thus be given below.

Clot formation
Blood clots were made using fresh venous blood from seven healthy volunteers not receiving anticoagulation treatment and with no history of coagulation disturbances. After collection the blood was immediately transferred to a Teflon-coated bottle. The
collected blood was then anticoagulated using citrate-phosphate-dextrose adenine (CPDA). Each blood clot was made from 1.4 ml CPDA-anticoagulated blood to which 0.025 mmol CaCl\(_2\) had been added to induce coagulation. The blood was then left to coagulate around a wool yarn (100 m/54 g, Peer Gynt, Sandnes Uldvarefabrik A/S, 4300 Sandnes, Norway) in a plastic test-tube for one hour [17, 18, 22].

**Determination of clot lysis**
Following one hour of coagulation, the clot was carefully extracted together with the wool yarn and mounted in a plastic frame that was lowered into a clot container with 160 ml r-PA or SK mixed NaCl solution [17, 18, 22].

To evaluate clot lysis 1 ml samples of the thrombolytic solution were taken from the clot container every 20 minutes during one hour (20, 40 and 60 min) to estimate the haemoglobin (Hb) leakage from the clot. The sample was added to 4 ml of Drabkins solution and the Hb content (mg) was measured by spectrophotometer at 540 nm, as described elsewhere [23]. To determine clot lysis the total Hb content (mg) of 1.4 ml anticoagulated blood mixed with 160 ml NaCl solution was measured (corresponding to a fully lysed clot) for each volunteer’s blood pool, the Hb content (mg) was then analysed as above. To determine clot lysis (% lysis) for each experimental clot, the loss of Hb (mg) in each clot was divided by the Hb content (mg) of the total Hb content of the 1.4 ml anticoagulated blood (fully lysed clot), resulting in an estimation of percentage clot lysis ((experimental clot Hb (mg)/ fully lysed clot Hb (mg)) \cdot 100).

**Thrombolytic drugs**
Two thrombolytic drugs were used in the present study, 0.25 U of r-PA (Rapilysin 10U\(^\text{®}\), Roche Registration Ltd, Hertfordshire, Great Britain) was mixed in 160 ml 0.9
% NaCl solution resulting in a concentration of 0.001562 U/ml. The other was SK, (Streptase®, 1.5 million international units, Hoechst Marion Roussel AB, Stockholm, Sweden), and 3600 IU mixed in 160 0.9 % NaCl solution with a resulting concentration of 22.5 IU/ml [22].

**Ultrasound exposure**
Continuous-wave US emitted by an unfocused piezoelectric transducer (CERAM AB, Lund, Sweden) with a resonance frequency of 1 MHz and diameter 16 mm was used in all experiments. The transducer was excited by an electronic system consisting of a function generator (HP 3314A, Hewlett-Packard, Washington, USA) and an RF power amplifier (ENI 240L, ENI, Rochester, New York, USA).

The effect of one hour of 1 MHz CW-US exposure, at intensities 0.0125, 0.025, 0.05, 0.1, 0.15, 0.3, 0.6, 0.9 and 1.2 W/cm² on clot lysis induced by either SK or r-PA was evaluated. For each thrombolytic drug and intensity where six interventional (US-exposed) clots submerged in thrombolytic solution and exposed to CW-US while six control clots (also submerged in thrombolytic solution) were left unexposed to US.

**Statistical analysis**
Wilcoxon’s signed rank test, was used to assess differences between interventional and control clots at each intensity, following 20, 40 and 60 min of exposure. In all statistical comparisons, P-values below 0.05 were considered significant.
**Ethical considerations**
The experiments described in the present study were conducted with the consent of each participant, and were approved by the Regional Ethical Review Board in Lund (approval: 879/2004).

**Results**

**Streptokinase treated clots**
Statistically significant decreases in clot lysis were seen at 0.9 W/cm$^2$ following 20 min (-2%), 40 min (-2%) and 60 min (-4%) CW-US exposure of SK-treated clots (P < 0.05). At an intensity of 1.2 W/cm$^2$ the decrease in clot lysis following 20 min of CW-US exposure was 3% and following 40 and 60 min of CW-US exposure 3 and 8%, respectively (P < 0.05 in all cases) (see Table 1 and Figure 1). No increase in clot lysis was seen in clots treated with SK and CW-US at any time or intensity of US exposure.

**Reteplase treated clots**
In the experiments using r-PA, statistically significant increases (P < 0.05) in clot lysis were seen at intensities of 0.05 W/cm$^2$ (3 %, P < 0.05), and at 0.3 W/cm$^2$ following 40 and 60 min of CW-US exposure (1 %, P < 0.05 and 8 %, P < 0.05) and at 0.6 W/cm$^2$ following 40 and 60 min (5 %, P = 0.03 and 8 %, P < 0.05) of CW-US exposure. Increased clot lysis was seen at all times following CW-US exposure at intensities of 0.9 and 1.2 W/cm$^2$ (0.9 W/cm$^2$: 20 min: 3 %, P < 0.05, 40 min: 4 %, P < 0.05, 60 min: 7 %, P < 0.05 and at 1.2 W/cm$^2$: 20 min: 8 %, P < 0.05, 40 min: 15 %, P < 0.05, 60 min: 10 %, P < 0.05). No significant decrease in lysis was seen at any time or US intensity in clots treated with r-PA (see Table 1 and Figure 1).
Discussion
The use of high frequency US to enhance thrombolysis during the treatment of stroke has shown promising results [1, 3]. However, clot lysis levels in vivo have not been in the same as those reported in vitro [7, 9, 10]. This may well be due to the attenuation of intensity as US passes through the temporal bone during high-frequency US exposure [11]. In the CLOTBUST trial, an intensity of 0.75 W/cm² was used [1], which would result in intensities between 0.01 and 0.06 W/cm² (following attenuation) reaching the obstructed vessel and the thrombus. We previously observed no enhanced fibrinolytic effects during pulsed-wave US exposure of r-PA-treated clots in this range of intensities [18], however effects were seen in the small intensity range between 0.125 and 0.25 W/cm². In the present study, using CW-US exposure, a statistically significant increase in lysis of r-PA treated clots was seen at low intensity (0.05 W/cm², 3 % increase, P = < 0.05). Thus, applying high-frequency CW-US to r-PA treated stroke patients may improve clinical results. However, different frequencies were used in the present study (1 MHz) and in earlier clinical studies (2 MHz) [1, 3], and the results should therefore be compared with care. Another explanation could be that the number of patients included in the CLOTBUST-trial was to small to achieve statistical significance, this despite efforts to include a sufficient number of patients [24].

In the present study, increasing enhancement of the clot lysis was seen in the experiments on r-PA treated clots (intensities ≥ 0.3 W/cm²), which are intensities higher than can be expected after passing through the skull bone [11]. In the present experiments on r-PA treated clots, enhancement of lysis was at lower intensities compared to our earlier study using pulsed-wave US exposure [18]. However, it would be difficult reach such levels of intensity with either pulsed or CW-US, due to
the high attenuation of the skull bone and considering the limited levels of output intensity recommended in transcranial Doppler US [25].

In the present study no enhanced effects of SK were seen at any intensity level used. This is a contradictory result when compared to results from studies with pulsed-wave US exposure [16, 17]. Thus, the mechanisms by which US enhance clot lysis might vary between CW-US exposure and pulsed-wave US exposure. This study shows decreased effects of clot lysis at intensities \( \geq 0.9 \, \text{W/cm}^2 \), a finding that has been seen earlier during pulsed-wave US exposure [16, 17], however at higher intensities (\( \geq 2 \, \text{W/cm}^2 \)). This might indicate that also duty cycle is an important factor influencing the results in US-enhanced clot lysis, not only intensity. Streptokinase is still considered by some to be a useful thrombolytic drug in the clinical setting [26, 27]. However, it does not appear to be suitable based on the decrease in effects induced by US exposure, according to the results of the present and earlier studies [16, 17]. Streptokinase has also been shown to be associated with a higher risk of intracranial bleeding than other thrombolytic drugs, and is therefore not recommended for clinical use in the treatment for stroke [28].

We have previously demonstrated a direct effect on the thrombolytic substance during exposure with pulsed-wave US exposure, effects associated with both decreased and increased effects on clot lysis [17, 18]. This was not examined in the present study, and we therefore do not know whether this effect exist when using CW-US. A recent study failed to reveal any changes in enzymatic activity of both SK and r-PA following US exposure [29]. Direct effects on the molecules of thrombolytic drug following US exposure must therefore be investigated in future studies.
Conclusions
Increasing intensities of CW-US exposure resulted in increased clot lysis of r-PA-treated blood clots, but decreased clot lysis of SK-treated clots. Continuous-wave US may thus be useful in US-enhanced clot lysis during stroke treatment with r-PA.

Competing interests
None.

Authors' contributions
Author BMH designed the investigation, performed the experiments, the statistical analysis and interpretation of the results, as well as the preparation of the manuscript.

Author JC assisted with the statistical analysis and the preparation of the manuscript.

Author AR supervised and designed the investigation as well as participated in the preparation of the manuscript.

All authors read and approved the final manuscript.
References


18. Hardig BM, Persson HW, Olsson SB: Direct action on the molecule is one of several mechanisms by which ultrasound enhances the fibrinolytic effects of reteplase. Blood Coagul Fibrinolysis 2006, 17:105-112.


Figures

**Figure 1 - Difference in clot lysis following continuous-wave ultrasound exposure**

Difference in clot lysis following 20, 40 and 60 min of continuous-wave ultrasound exposure at different intensities, presented as the difference between ultrasound-exposed clots and control clots: □ = clots exposed continuous-wave ultrasound and streptokinase, ■ = clots exposed to continuous-wave ultrasound and reteplase.

Wilcoxon’s signed rank test was used to assess statistical differences, * = P < 0.05.

Tables

**Table 1 - Difference in clot lysis following continuous-wave ultrasound exposure**

Clot lysis (%) of clots exposed to streptokinase concomitantly with continuous-wave ultrasound at different intensities for one hour (US+SK) and clots exposed to streptokinase alone (Controls) and in clots exposed to reteplase concomitantly with continuous-wave ultrasound at different intensities for one hour (US-r-PA) and reteplase alone (Controls). Results are presented as medians and 5\(^{th}\) – 95\(^{th}\) percentiles.

Wilcoxon’s signed rank test was used to assess statistical difference.
Figure 1
Additional files provided with this submission:

Additional file 1: table_01_clotlysis_us.doc, 88K
http://www.biomedcentral.com/imedia/1878975445187507/supp1.doc