The Novel Anti-Thrombotic Drug with No-Bleeding Excess

Mark IM Noble1*, Isobel Ford1, Gary Cameron1 and Angela Drake-Holland2
1Department of Medicine and Therapeutics, University of Aberdeen, Scotland
2School of Pharmacy, Robert Gordon University, Scotland
Submission: May 10, 2017; Published: June 30, 2017
*Corresponding author: Mark IM Noble, Department of Medicine and Therapeutics, University of Aberdeen, Scotland, Email: mimnoble@abdn.ac.uk

Abstract

Background and purpose: All anti-thrombotic drugs presently available cause increased bleeding, a most undesirable effect. Our aim was to test a novel anti-thrombotic drug [Th001] which, theoretically should not suffer from this disadvantage.

Experimental approach: The trial design was a double-blind, placebo controlled, paired crossover, with 99% power for showing a difference in skin bleeding time. n=48 Caucasian patients, age 69.0±6.6 years, 38 male, 10 female, with stable atherothrombotic disease and other stable chronic diseases, treated with aspirin and statin. There was no effect on bleeding time (p = 0.9729) nor on tests of haemostasis, (Ultegra point of care aggregation test and flow cytometry). The mean platelet count rose from 234.43±67.264 to 246.43±84.14x10-9 per litre, p=0.024.

Conclusion: Th001 therapy appears to be a safe method for potential treatment and prevention of occlusive arterial thrombosis with no risk of increased bleeding.
Keywords: Arterial thrombus growth inhibition; Coronary thrombosis; Thrombotic stroke; Peripheral arterial disease; Normal haemostasis

Introduction

This clinical study was performed in 2006-2007, on a drug for which the preclinical results were published, when it was entitled ICI 170809 [1]. Publication submission for this clinical study has been delayed for a variety of reasons, viz:

I. Failure to attract sufficient academic grants to develop the drug for clinical use.

II. Death of Professor Gabrielle Hawksworth, who was the senior pharmacologist on the study. This publication is a memorial to her.

III. Lack of effective intellectual property for an old drug that lost its molecular patent in 2003.

IV. Delay in forming a company that sought such intellectual property, which was a long process ending in the grant of a US patent of use in December 2015.

V. In spite of favourable reception from several referees, journals have declined publication in view of the age of the data. Contrary to that view is the opinion of [2], in the book ‘Bad Pharma’, namely the insistence that all results of drug trials should be published, regardless of age.

Serotonin is involved in platelet-rich thrombus growth in diseased, stenosed arteries (with increased shear rates and endothelial damage). Serotonin is not involved in the initial platelet activation, aggregation and fibrinogen binding to form a haemostatic layer when normal tissue is damaged; such tissue contains no serotonin. Conventional drug discovery for anti platelet drugs requires that there should be a measurable inhibitory effect upon conventional platelet function tests which all measure haemostatic platelet function; such drugs will inevitably cause bleeding. Drug discovery for an anti-thrombotic agent that does not cause bleeding should not have these effects. Arterial thrombosis tends to occur at arterial narrowings, pointing to the probability that the convective acceleration, generating increased shear stress, is the predominant stimulus to platelet-rich thrombus growth at these sites. When platelet deposition occurs in an arterial stenosis, serotonin is released from the dense granules of the platelets and activates more
platelets through the 5HT2A receptors, leading to a platelet-to-platelet positive feedback reaction (snowball)-the platelet-rich thrombus growth. Therefore, platelet-rich thrombus growth can be differentially blocked by antagonising the platelet serotonin receptor, of the 5HT2A subtype, leaving haemostatic function intact. That this occurs with the 5HT2A antagonist Th001 (formerly ICI 170809) is known from the work of McAuliffe et al. [1]. This was shown in the “gold standard” method for this purpose—the Folts model of the anaesthetised dog with damaged and stenosed coronary artery [3]. These animals, which are studied during a thoracotomy, show no increased surgical bleeding upon administration of the ICI 170809, now known as Th001 (Arteclere™). There is thrombus dispersal that is not due to a direct action of Th001, but to the effect of natural thrombolysis when thrombus growth is abolished. Although this measurement cannot be performed in man, there is no reason to think that there is a great difference between man and dog. The Folts model has always reliably predicted the efficacy of anti-platelet drugs in man [3].

We report the results of an academic study funded by grant to the University of Aberdeen, carried out on consenting patients at Aberdeen Royal Infirmary, monitored by the Research and Development department, and approved by MHRA. The patients were stable with controlled atherothrombotic arterial disease, either coronary or peripheral, attending as out-patients. The results failed to demonstrate any significant effects of Th001 upon a number of indices of haemostasis.

**Methods**

**Patients**

The study, including patient information sheet, invitation letter to patients, informed consent form, and GP information letter were approved by Grampian Region Ethics Committee, conforming to BJR guidelines. Patients with inclusion criteria were invited to participate by letter. This was a real patient situation because of minimal exclusion criteria. Inclusion patients previously diagnosed with either coronary arterial disease, defined as ≥70% stenosis on angiography or peripheral arterial disease, with an/diage/brachial pressure ratio <0.9 these patients were treated with aspirin and a statin. Exclusions-taking warfarin or clopidogrel, leg ulceration and rest pain, females of child bearing age, and patients receiving serotonin re-uptake inhibitors. 46 patients were recruited with ischaemic heart disease plus 10 patients with peripheral vascular disease. A total of 48 patients, after exclusions completed the study. We report the completed studies on 48 Caucasian patients, age 69±6 years, 38 male, 10 female, with stable athero-thrombotic disease treated with aspirin and statin. The entire study was carried out from 2006 to 2007.

The co-morbidities included hyperlipidaemia (treated) 23, hypertension(treated) 20, diabetes mellitus 11, renal disease 9, hepatic insufficiency 8, left ventricular dysfunction 7, hypothyroidism 6, chronic obstructive pulmonary disease 5, iron deficiency anaemia 5, diverticular disease; duodenal disease 4, hearing loss 4. Other diseases in fewer numbers of patients each were: haematuria, prostatic disease, urinary incontinence, back pain, macrocytic anaemia, asthma, glaucoma, haemorrhoids, hiatus hernia, impotence, Meniere’s disease, osteoporosis, retinopathy, sciatica, ankylosing spondylitis, anxiety and depression, aortic aneurysm, aortic stenosis, cerebral spondylosis, Charcot foot, Crohn’s disease and macular degeneration. These diseases were considered to be stable by the patients’ physicians. The past histories of these patients included bacterial meningitis, basal cell carcinoma, cervical polypl, colonic polypl, CVA, pneumonia, pulmonary tuberculosis, numerous and various operations.

The study design was double-blind, placebo controlled cross over. Subjects attended the Cardiology Research suite for five visits: control pre-treatment baseline, on course 1 for 2 weeks, washout for 2 weeks, on course 2 for 2 weeks, washout for 2 weeks. Course 1 was randomized to placebo or Th001 10mg bd and reversed for course 2. The randomization and preparation of courses was carried out by an independent agency at Ninewells Hospital, Dundee. All participants in the study -patients, staff and investigators were blinded to the randomization code. At each visit, physical measurements were: clinical examination plus electrocardiogram (ECG), including heart rate, QT interval and QT interval corrected for heart rate (QTc), followed by:

**Venepuncture:** Venous blood was collected into vacutainers by venepuncture with 19G butterfly and adaptor. The first 5ml of blood was used for full blood count, and subsequent samples collected in order for the platelet tests and others.

**Platelet aggregation tests:** Platelet aggregation was measured using the RPFA Ultegra [4], with Verify now cartridges for aspirin and GPIIB/IIIa (Accumetrics, San Diego, CA). This is a well-validated, point-of-care system which tests the ability of activated platelets in whole blood to bind fibrinogen-coated micro-beads contained within the cartridge test wells. Blood is stimulated with either thrombin-receptor activating peptide (TRAP) in the GPIIB/IIIa assay, or arachidonic acid (AA) in the aspirin assay. Results are reported as platelet aggregation units (PAU) for the GPIIB/IIIa assay and aspirin reaction units (ARU) for the aspirin assay (>550 ARU indicates a poor- or non-response to aspirin. Our inter-assay variation is 5.8% for RPFA-Aspirin and 5.0% for RPFA-GPIIB/IIIa.

**Platelet activation:** This was assessed as fibrinogen binding to resting and stimulated platelets by whole blood flow cytometry [5], as previously validated [6]. Briefly, citrate-anticoagulated blood was diluted ten-fold with HEPES-Mg buffer within 10 minutes of blood sampling. Diluted blood was incubated with FITC-conjugated rabbit anti-human fibrinogen antibody (Dako Cytomation, Denmark), for 20 minutes and the reaction was stopped by dilution with cold PBS. In order to

How to cite this article: Mark I, Isobel F, Gary C, Angela D H. The Novel Anti-Thrombotic Drug with No-Bleeding Excess. J Cardiol & Cardiovasc Ther 2017; 6(2): 555684. DOI: 10.19080/JOCCT.2017.06.555684.
Skin bleeding time was performed, We tested variables from placebo and Samples were blinded and batch

The Novel Anti-Thrombotic Drug with No-Bleeding Excess. J Cardiol & Cardiovasc Ther

paper (Whatman No.1) was applied to the droplet at 30 second (Elitech, Hertfordshire), and a stopwatch was started. Filter

40mmHg. One incision was made with Surgicutt bleeding device sphygmomanometer cuff was inflated on the upper arm to

was shaved if necessary and cleaned with alcohol swab. A contra-laterally from venepuncture. A suitable area of forearm

still detectable during course 2 (placebo) period.

Plasma Th001 assay: Samples were blinded and batch processed, after the end of the study, by a staff scientist, in order

that the code could not be broken. Plasma was prepared by centrifugation of citrate-anticoagulated blood at 2500×g for 10

minutes, within 1 hour of venepuncture, frozen, stored at -80 °C until assay. Plasma Th001 concentrations were determined using a thermos Surveyor-TSQ Quantum LC-MS/MS system (Hemel Hempstead, UK). Briefly, 50 µl of plasma were precipitated by vortex mixing with 200 µl of acetonitrile containing 100 ng/ml imipramine as the internal standard and centrifuged at 10,000 rpm for five minutes. 5.0 µl of the supernatant was injected onto the chromatograph. Imipramine and Th001 (formerly ICI 178009 and obtained from ICI Pharmaceuticals) were resolved on a Nucleosil 5u ODS column (150 ×6.6 mm) with a mobile phase of 20% 0.1% triethylamine, pH 4.0 and 80% acetonitrile at a flow rate of 1 ml/min, which was split to allow 200 µl/min through the mass spectrometer. The first four minutes after injection were diverted to waste. Both compounds were analysed using an electrospray source in positive ion mode and quantification performed on the SRM transition fragments m/z 337.1-m/z 292.1 for Th001 and m/z 281.1-m/z 86.1 for imipramine. Calibration standards and quality control samples were prepared by spiking blank plasma with known concentrations of Th001 and processing as described for the patient samples. Calibration curves were linear up to 1000 ng/ml with a lower limit of quantification of 1 ng/ml.

Compliance: Compliance was determined after the end of the study by examination of the plasma Th001 concentrations. Plasma Th001 concentration assays showed 100% compliance, but the concentrations during Th001 administration were 89.1±59.0 ng/ml. In 6 patients who received Th001 as course 1 and achieved the highest peak concentrations, a little Th001 was still detectable during course 2 (placebo) period.

Skin bleeding time: Skin bleeding time was performed, contra-laterally from venepuncture. A suitable area of forearm was shaved if necessary and cleaned with alcohol swab. A sphygmomanometer cuff was inflated on the upper arm to 40 mmHg. One incision was made with Surgicutt bleeding device (Elitech, Hertfordshire), and a stopwatch was started. Filter paper (Whatman No.1) was applied to the droplet at 30 second intervals, avoiding touching the skin, until bleeding stopped. Bleeding time was recorded to the nearest 30 seconds. These measurements were made by a trained nurse, blinded to the phase of the treatments.

Other observations: Measurements made by Aberdeen Royal Infirmary (ARI) routine full blood count, Clinical Haematology Laboratory; urea and electrolytes (U&Es), creatinine, Liver Function tests (LFTs) and blood glucose, Clinical Biochemistry Laboratory. Prothrombin fragments F1+2 by the University Platelet Laboratory.

Statistical analysis: We tested variables from placebo and Th001 patient data for Gaussian normality distribution by the Kolmogorov and Smirnov method. Normally distributed data sets were then analysed by paired t test, and non-normally distributed data were analysed by Wilcoxon paired signed rank test. Two weekly interval measurements of platelet count were analysed by repeated measures ANOVA. In Stat software (Graph Pad, San Diego) was used on a MacIntosh Apple computer (iMac).

Results

Th001 assays

We were aiming at 100 µg/kg body wt. As there are 70 ml blood per kg body wt, this is equivalent to 7 µg/litre, i.e., 7 ng/ml. During Th001 administration, plasma Th001 concentration was 91.47±60.09 ng/ml. The levels achieved are an order of magnitude higher than planned.

i. In all patients during the Th001 period, Th001 was detected.

ii. In all patients taking placebo before Th001, no Th001 was detected.

iii. In 30 patients, Th001 was detected in the washout period following the Th001 period.

iv. In 6 of these patients who took Th001 before placebo, Th001 was detected during the placebo period, i.e., carried over from the Th001 period.

v. In 4 of these patients, Th001 was detected in the washout period following the placebo period.

The statistically analysed groups of variables relevant to haemostasis are presented in (Table 1). There was a statistically significant increase in platelet count in those patients receiving Th001 first. The effect on platelet count is more clearly shown by repeated measures analysis of variance in the patients given Th001 first (Figure 1). The effect of Th001 upon biochemical variables is presented in (Table 2). All comparisons made by Wilcoxon. There were small but statistically significant rises of LFTs in patients taking Th001. With few exceptions the changes were within the normal range. In patients whose LFTs were elevated in the control period, a significant further rise on Th001 was found only for gamma GT. The increase in LFTs would not have been detected but for the paired design; the increases
were almost all small and within the normal range. This effect may not be detectable at a lower dose, e.g., 5 mg od. Prothrombin fragments F1+2 were mostly in the normal range and showed a non-significant decrease in patients to whom Th001 was administered. The results of standard haematology tests are presented in (Table 3).

**Table 1: Bleeding Time, Platelet Count, Flow Cytometry.**

| Variable                                      | Probability of No Difference | Placebo (Mean ±SD) | Th001 (Mean±SD) | Test Used    |
|-----------------------------------------------|------------------------------|--------------------|-----------------|--------------|
| Bleeding Time                                 | NS                           | 310.4±76.6 sec     | 311.7±82.09     | paired t test|
| Platelet Count                                | 0.024                        | 234.7±67.3x109/l   | 246.4±84.1x109/l| paired t test|

**Platelet Function**

| Variable                                      | Probability of No Difference | Placebo | Th001 | Test Used |
|-----------------------------------------------|------------------------------|--------|-------|-----------|
| Ultegra ASA aspirin reaction units            | NS                           | 441.4±63.7 ARU | 440.1±54.4 ARU| Wilcoxon   |
| Ultegra IIb/IIIa platelet aggregation units   | NS                           | 209.7±67.1 PAU  | 211.6±74.4 PAU | paired t test|
| Flow cytometry: resting fibrinogen (%bound)  | NS                           | 3.10±3.60%      | 2.78±3.41%     | Wilcoxon   |
| Flow cytometry + 10 µM ADP (%fibrinogen bound with ADP stimulation) | NS | 55.59±19.12% | 54.98±18.69%| paired t test|
| Flow cytometry + 10 µM ADP (%fibrinogen bound with serotonin stimulation) | NS | 36.14±18.55% | 35.57±17.43%| Wilcoxon   |
| Flow cytometry + 10 µM ADP + 10 µM 5HT (%fibrinogen bound with ADP + serotonin stimulation) | NS | 38.15±19.04% | 43.11±21.83%| paired t test|

**Table 2: Biochemical variables.**

| Variable               | p value for Th001 vs Placebo | Placebo (Mean ±SD) | Th001 (Mean±SD) | Test Used |
|------------------------|------------------------------|--------------------|-----------------|-----------|
| Bilirubin              | NS                           | 9.313±5.007µM      | 9.021±5.393µM   |           |
| Alkaline phosphatase   | 0.0082                       | 78.77±34.24U/l     | 82.34±36.49U/l  |           |
| Alanine amino transferase | 0.0021                       | 26.77±11.41U/l     | 30.21±15.86U/l  |           |
| Gamma GT               | 0.0216                       | 37.00±3.60U/l      | 42.75±3.85U/l   |           |
| Glucose                | NS                           | 6.93±2.76mM        | 7.14±3.12mM     |           |
| Urea                   | NS                           | 6.31±1.77          | 6.58±2.05mM     |           |
| Creatinine             | NS                           | 100.5±19.8mM       | 102.3±20.3mM    |           |
| Sodium                 | NS                           | 139.5±2.4mM        | 139.8±2.2mM     |           |
| Potassium              | NS                           | 4.28±0.39mM        | 4.22±0.38mM     |           |
| Chloride               | NS                           | 105.5±2.4mM        | 106.1±2.7mM     |           |

**Table 3: Standard Haematology.**

| Variable               | Probability of No Difference | Placebo (Mean ±SD) | Th001 (Mean±SD) | Test Used |
|------------------------|------------------------------|--------------------|-----------------|-----------|
| Haemoglobin            | NS                           | 139.5±11.8 g/l     | 138±12.1g/l     |           |
| Red Blood Cell Count   | NS                           | 4.566±0.462        | 4.534±0.414     |           |
| White Blood Cell count | NS                           | 7.687±2.271        | 7.843±2.058     |           |

**Figure 1:** Platelet counts in patients receiving Th001 at a dose of 10 mg twice daily first. Mean±standard deviations at control, after 2 weeks of Th001 treatment, after 2 weeks of washout and after 2 weeks of placebo. p=probability of no difference analyzed by repeated measures ANOVA.

**ECG and adverse events**

There were no statistically significant differences in QT interval or corrected QT interval (QTc) in any of the electrocardiograms. Adverse events were scaled as 0 = denied, 1 =mild, 2 =moderate, 3 =severe. They were present in 17 patients at scale 1, the remainder at scale 0. The difference between adverse events in the placebo and Th001 periods was not statistically significant (p =0.59). No significant drug interactions were found. However, one patient whose control heart rate was 48/min experienced a fall of heart rate to 37/min while taking Th001 and required reduction of his atenolol dose. Two patients taking bisoprolol had slight increases in heart rate. The remaining patients on β-adrenoreceptor antagonists experienced variable heart rate responses, which were significantly reduced in this...
sub-group. There was no significant change in heart rate in patients not taking β-adrenoreceptor antagonists.

**Discussion**

Efficacy of the inhibition of arterial thrombosis by serotonin 5HT2A antagonists is attested by numerous animals studies of the effect of 5HT2A antagonists [7-17]. Macroaggregate platelet-rich aggregate growth is inhibited by ICI 170809 in hirudinised human blood by single platelet counting [18].

The important aspects of the present study are (1) lack of risk of bleeding and (2) lack of adverse events, both indicating that Th001 is a safe drug for the treatment and prevention of platelet-rich arterial occlusive thrombosis, assuming that efficacy for that effect can be shown in man. The small effect on liver function (similar to that found in a larger cohort by ICI) would not have been found without the paired design, and may disappear when the dose is reduced to 5mg od from the 10mg bd used here. The choice of 10mg bd was fortunate in that the resulting plasma concentrations were far above any likely levels in future studies with lower doses; this reinforces the value of the safety data. An effective dose may be as low as 5mg od. Also, with lower doses, patients are less likely to experience β-adrenoreceptor antagonist induced bradycardia below 60/ min requiring the dose of β-adrenoreceptor antagonist to be reduced.

The dose of 10mg bd was adopted upon the advice of one of the co-authors of the paper by [9]. When the plasma samples were assayed after the end of the study, it was found that the values were approximately four times those that were maximally effective in the study by [1], in dogs. This miscalculation led us to conclude that it enhanced the value of the safety data as no significant adverse event were recorded with this excessive dose apart from the effect on liver function tests. It is a little surprising to see a persistence of the drug after 2 weeks at a dose of 10mg. The fact that this is seen in only some individuals points to variations in drug metabolism (e.g. by cyp polymorphisms), or excretion. Full details of these considerations are available in the ICI (now Thromboserin Ltd) archive.

The assessment of platelet function lacks specific analysis of pathways that might be affected by 5HT in vivo. 5HT is well known to augment the platelet response to other agonists, while not acting as a primary agonist at physiological concentrations. We saw very little response of the platelets to 5HT alone. Since it cannot be acting as a primary agonist at physiological concentrations. We hesitate to interpret the increase in platelet count in patients receiving Th001 in the first treatment period. Our speculation that this could be accounted for as a reduction in platelet consumption would need to be tested and is best dismissed for the time being as an speculation influenced by bias (see conflict of interest). The lack of bleeding risk indicates that Th001 would be a potential safe therapy (if efficacy can be proven in patients) for the prevention of arterial thrombotic occlusion around the time of surgical operations, when other anti-thrombotic treatments are contra-indicated because of surgical bleeding. Our patients were stable, and therefore had little, if any growing thrombi at arterial stenoses. Such growing thrombi are characteristic of acute thrombus related clinical conditions in arterial disease.

**Conclusion**

We conclude that the absence of any pro-haemorrhagic effect of Th001, and the excellent safety profile thus far, may make this a most suitable anti-thrombotic agent for further development.

**Acknowledgement**

The senior pharmacologist supervising this study, Professor Gabrielle Hawsworth, is now deceased. We dedicated this write-up to her memory. Many thanks are due to clinical staff in the cardiology research room, and technicians in the University laboratories, for their work and support. The funding consisted of academic grants to the University of Aberdeen from the Coronary Thrombosis Trust, Enterprise Scotland Grampian and Tenovus Scotland.

**Conflicts of Interest**

There were no conflicts of interest at the time of the study (2006-2007) but subsequently Noble and Drake-Holland became share holders in Thromboserin Ltd who hold the patents of use for Th001, formerly ICI 170809, the drug used in this academic study.

**References**

1. McAuliffe SJG, Snow HM, Cox B, Smith CC, Noble MIM (1993) Interaction between the effect of 5-hydroxytryptamine and adrenaline on the growth of platelet thrombi in the coronary artery of the anaesthetised dog. Br J Pharmacol 109(2): 405-410.
2. Goldacre B (2012) Bad Pharma. Harper & Collins, USA, ISBN: 86010095446.
3. Torr S, Noble MIM, Folts JD (1990) Inhibition of acute platelet thrombosis formation in stenosed canine coronary arteries by the
specific serotonin 5HT2 receptor antagonist ritanserin. Cardiovasc Res 24(6): 465-470.

4. Husseini HM, Emiru T, Georgiadis AL, Qureshi AI (2013) Assessment of platelet inhibition by point-of-care testing in neuroendovascular procedures. Am J Neuroradiol 34(4): 700-706.

5. Goodall AH, Appleby J (2004) Flow-Cytometric Analysis of Platelet-Membrane Glycoprotein Expression and Platelet Activation. Methods Mol Biol 272: 225-253.

6. Cassar K, Bachoo P, Ford I, Greaves M, Brittenden J (2003) Platelet activation is increased in peripheral arterial disease. J Vasc Surg 38(1): 99-103.

7. Ashton JH, Benedict CR, Fitzgerald C, Rabeja S, Taylor A, et al. (1986) Serotonin as a mediator of cyclical flow variation in stenosed canine coronary arteries. Circulation 73(3): 572-578.

8. Belcher PR, Drake-Holland AJ, Hynd J, Noble MIM (1992) Dispersion of coronary artery thrombi by antagonism of platelet serotonin receptor in the dog. Cardiovasc Res 26(3): 292-296.

9. Milkon DS, Jessop CL, Swaibland A, Haworth S, Rushston A, et al. (1992) The effect of a selective 5HT2 receptor antagonist (ICI 170,809) on platelet aggregation and pupillary responses in healthy volunteers. Br J Clin Pharmacol 33(3): 281-288.

10. Kirchengast M, Rubsamen K, Lehmann HD (1993) Inhibition by the combined Ca2+ and 5-HT2 receptor antagonist nesopamil (LU 49938) of intracoronary thrombus formation in a canine model of arterial stenosis and intimal damage. J Cardiovasc Pharmacol 22: 687-694.

11. Hsieh CP, Sakai K, Bruns GC, Dage RC (1994) Effects of MDL 28,133A, a 5-HT2 receptor antagonist, on platelet aggregation and coronary thrombosis in dogs. J Cardiovasc Pharmacol 24(5): 761-772.

12. Pawlak D, Pawlak K, Chablienska E, Malyszko J, Takada A, et al. (1998) A potent 5HT receptor (5-HT2A) antagonist, BV-7028, delays arterial thrombosis development in rats. Thromb Res 90(6): 259-270.

13. Berry CN, Lornin J, Lockett S, Delahaye M, Lale A, et al. (2001) Antiplatelet and antithrombotic activity of S65.0472, a mixed 5HT1B/5-HT2A receptor antagonist. Thromb Haemost 85(3): 521-528.

14. Kihara H, Koganei H, Hirose K, Yamamoto H, Yoshimoto R (2001) Antithrombotic activity of AT-1015, a potent 5-HT2A receptor antagonist, in rat arterial thrombosis model and its effect on bleeding time. Eur J Pharmacology 21: 157-162.

15. Adams JW, Ramirez J, Ortuno D, Shi Y, Thomsen W, et al. (2008) Anti-thrombotic and vascular effects of AR246686, a novel 5-HT2A receptor antagonist. Eur J Pharmacol 586(1-3): 234-243.

16. Przyklenk K, Frelinger Al, Linden D, Whittaker P, Li Y, et al. (2010) Targeted inhibition of the serotonin 5HT2A receptor improves coronary patency in an in vivo model of recurrent thrombosis. J Thromb Haemost 8(2): 331-340.

17. Lin OA, Karim ZA, Vemana HP, Espinosa EV, Khasawneh FT (2014) The antidepressant 5-HT2A receptor antagonists pizatifen and cyproheptadine inhibit serotonin-enhanced platelet function. PLoS One 23: e87026.

18. Menys VC (1993) Collagen induced human platelet aggregation: serotonin receptor antagonism retards aggregate growth in vitro. Cardiovasc Res 27(11): 1916-1919.