Lessons from the trials

The Prospective Randomized On-X Valve Anticoagulation Clinical Trial (PROACT): Lower is better, but is it good enough?

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ABSTRACT

Due to their durability, mechanical prostheses are frequently used for aortic valve replacement (AVR) in young adults. However, these valves are thrombogenic and require lifelong anticoagulation. Over the last few decades, efforts have been made towards the lowering of INR targets in an effort to reduce bleeding events without influencing the thromboembolic risk. The Prospective Randomized On-X Valve Anticoagulation Clinical Trial (PROACT) was designed to compare standard versus low anticoagulation targets in high-risk patients undergoing mechanical AVR with the ON-X prosthesis.

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INTRODUCTION

The ideal aortic valve substitute for young adults remains elusive. While bioprostheses avoid long-term anticoagulation, durability is a major concern in young adults because of early structural valve degeneration.\(^1,2\) Due to their durability, mechanical prostheses are more frequently used in young adults. However, these valves are thrombogenic, requiring lifelong anticoagulation which is mostly responsible for valve-related morbidity and mortality following mechanical AVR.\(^2-4\) This partially explains the increased observed mortality in young adults undergoing mechanical AVR in comparison with an age- and gender-matched general population.\(^2\) Therefore, reducing anticoagulation-related complications are major ongoing research objectives to improve survival and quality of life for this patient population. Over the last decades, efforts have been made towards lowering of INR targets, which resulted in a reduction of bleeding-event rates without influencing the risk of thromboembolic (TE) events.\(^5,6\) Additionally, improvements in mechanical prosthesis engineering have led to a substantial reduction in the thrombogenicity of these devices and subsequently a decrease in the incidence of valve-related complications.\(^7\) However, the impact of new bileaflet mechanical prostheses on the safety of lowering anticoagulation targets remains uncertain. The Prospective Randomized On-X Valve Anticoagulation Clinical Trial (PROACT) aimed to evaluate the safety of lower anticoagulation (target INR 1.5–2.0) after implantation of the ON-X prosthesis (Figure 1) in patients at high risk of thromboembolic events.\(^8\)

THE STUDY

The PROACT trial was a multicenter unblended controlled randomized study published in 2014 in *the Journal of Thoracic and Cardiovascular Surgery*.\(^8\) The trial was designed to compare standard versus lower anticoagulation therapy in patients at high risk of TE events, undergoing mechanical AVR with the ON-X prosthesis, including patients with chronic atrial fibrillation, left ventricular ejection fraction <30%, enlarged left atrium (50 mm), hypercoagulability or resistance to anti-platelet therapy and women receiving estrogen therapy. Ninety days after surgery, 425 patients were enrolled and 375 patients meeting the inclusion criteria were finally randomized in one of two study arms: (1) a targeted INR between 1.5 and 2 (test group) and (2) a targeted INR between 2 and 3 (control group). Concomitantly, all patients received aspirin 80 mg and were monitored using home INR testing. Any patient who experienced a TE event in the study group was
Table 1  Events rate comparison between patients in the test group and the standard INR group in the PROACT trial (adapted from Puskas et al.\\(^8\)).

| Event Type                        | Test group (%/pt-yr) | Control group (%/pt-yr) | \(P\) value |
|-----------------------------------|----------------------|--------------------------|-------------|
| Bleeding and TE events            | 5.63                 | 8.47                     | 0.046       |
| Major events (major bleeding, TE and thrombosis) | 4.44                 | 5.16                     | 0.539       |
| Major bleeding events             | 1.48                 | 3.31                     | 0.032       |
| Minor bleeding events             | 1.18                 | 3.31                     | 0.011       |
| All bleeding events               | 2.67                 | 6.62                     | <0.001      |
| TE and thrombosis events          | 2.96                 | 1.85                     | 0.178       |
| All cause mortality               | 1.48                 | 1.46                     | 0.968       |

Notes.
- TE, thromboembolic.

crossed over to the standard INR group, though they remained in the test group through an intention-to-treat analysis. No patients were allowed to cross over from the standard to the test group. The study was sponsored and funded by On-X Life Technologies (Austin, TX) and conducted under an investigational device exemption provided by the US Food and Drug Administration. Primary endpoints were rates of TE events, thrombosis, bleeding and all-cause mortality. An interim non-inferiority analysis was performed for a composite of all primary endpoints, with an absolute margin of 1.5%.

The mean follow-up was 3.8 years and was 98% complete. The mean age was 55 years. The two groups were comparable in terms of patient demographics, except for a marginally higher prevalence of preoperative atrial fibrillation in the control group (6% versus 2%, \(p = 0.06\)). The mean INR was 1.89 in the test group and 2.5 in the control group (\(p < 0.001\)).

In terms of findings, the composite of major and minor bleeding, TE and thrombosis events was significantly lower in the test group (5.63%/pt-yr in the test group versus 8.47%/pt-yr in the standard group, \(p = 0.046\)) (Table 1). Furthermore, the test group experienced a significantly lower rate of major bleeding events (1.48%/pt-yr in the test group versus 3.31%/pt-yr in the standard group, \(p = 0.032\)) and minor bleeding events (1.18%/pt-yr in the test group versus 3.31%/pt-yr in the standard group, \(p = 0.011\)) (Figure 2), with no differences in the rates of TE and thrombosis events (2.96%/pt-yr in the test group versus 1.85%/pt-yr in the standard group, \(p = 0.178\)) (Figure 3).

Interestingly, when comparing the composite of outcomes while excluding minor events, there was no statistically significant difference between the 2 arms (4.44%/pt-yr in the test group versus 5.16%/pt-yr in the standard group, \(p = 0.539\)).

DISCUSSION

The investigators concluded that, with the On-X prosthesis, a target INR between 1.5 and 2 translates into a lower incidence of bleeding events without a significant increase in TE events. This is consistent with recently published studies demonstrating the safety of a lower anticoagulation regimen in patients with mechanical AVR\\(^9–11\). While the authors state in the limitations that their results could not be extrapolated to other prostheses, the LOWERING-IT trial studied the impact of a lower anticoagulation regimen in patients with various mechanical aortic prostheses\\(^9\). In that study, patients were randomized to low dose INR (1.5–2.5) or standard INR (2–3) anticoagulation therapy. Similar to findings
from the PROACT trial, there were no differences in TE events (OR 0.33 [0.006–4.20], \( p = 0.6 \)) while there was a significant decrease in bleeding events in the low-dose group (OR 0.36 [0.11–0.99], \( p = 0.04 \)).

While the PROACT trial was generally well conducted, it raises several important points that warrant consideration. Firstly, the study sample size was calculated to demonstrate non-inferiority of a composite endpoint of bleeding, TE and thrombosis events, which was entirely driven by the reduction in bleeding events in the lower INR group as anticipated. Nevertheless, looking specifically at TE and thrombosis events, there was a 60% higher rate in the test group (2.96%/pt-yr versus 1.85%/pt-yr). Although this did not reach statistical significance (\( p = 0.178 \)), it may be attributed to a lack of statistical power due to sample size.

These findings require further analysis to ensure safety of aiming for lower INR targets. Secondly, as part of the study protocol, patients were provided a home INR monitoring kit and were closely followed up. This translated into the fact that out of 53,000 measurements, more than 60% of the measured INRs were within the desired range, with 96% of patients having at least one test per month. This correlates with previous studies where home INR monitoring was associated with better INR control, higher long-term survival and lower anticoagulation-related events in patients with mechanical prosthesis\(^{12-14}\). However, home INR testing is not widely adopted in the wider population because of availability and cost issues, and patient compliance with testing is overall lower in a real world setting\(^{15}\), which results in patients being off range a significant
portion of time. Consequently, results of this trial should be applied with caution in a daily practice because the lower range in the test group (INR 1.5) leaves little safety margin for being under-anticoagulated.

Thirdly, while there was a reduction in anticoagulation-related complications in the low INR group, rates of major bleeding and neurological events (1.48%/patient-year and 1.98%/patient-year, respectively) remain comparable to previously published cohort studies examining long-term outcomes following mechanical AVR. This suggests that these events may be under-reported and under-estimated in retrospective cohort studies. Importantly, the rates of major events (major bleeding, TE and thrombosis) were not different between the lower and standard anticoagulation groups in the PROACT trial. This has major implications in a young adult population with long anticipated life expectancy where the lifetime risk of experiencing one major anticoagulation-related event is a major consideration.

WHAT HAVE WE LEARNED?

The PROACT trial confirms that a lower anticoagulation target with the On-X prosthesis results in a reduced rate of bleeding and TE. This study represents a major step forward in outcomes improvement for patients with mechanical prostheses. In addition, this study reinforces the potential benefit of home INR monitoring on clinical outcomes in this patient population. Despite a lower INR target and a closer anticoagulation management, anticoagulation-related complications remain the main limitation following mechanical AVR. Furthermore, the real impact of reduced anticoagulation-related events on long-term survival and quality of life of patients, especially in the younger patient population, are yet to be determined. When applying it in clinical practice, this trial should be interpreted in light of recent data available for other valve substitutes. While bioprostheses do not require anticoagulation, durability is a major issue in young adults because of early structural valve degeneration and long-term survival remains suboptimal. Despite these concerns, their use in this population has gained adoption with the advent of transcatheter options for subsequent reintervention.

In contrast, the Ross procedure (pulmonary autograft replacement) alleviates the need for lifelong anticoagulation and is the only operation that guarantees long-term viability of the aortic valve substitute. In several recent reports, this has translated into long-term survival equivalent to the age- and gender-matched general population, a lower risk of valve-related complications and better quality of life than bioprosthetic and mechanical prostheses.

Recently, attempts have been made to broaden anticoagulation options in patients with mechanical prostheses. Novel oral anticoagulants (NOACs) are an alternative to warfarin that preclude the need for laboratory testing. Published in 2013, the REALIGN trial randomized patients to receive Dabigatran versus warfarin after mechanical valve replacement. The use of dabigatran was associated with increased rates of thromboembolic and bleeding events (Figure 4). However, the targeted serum levels of Dabigatran (≥50 ng per ml) was extrapolated from studies on stroke prevention in patients with atrial fibrillation and may be insufficient for prosthetic mechanical valves.

In addition, all major bleeding events were pericardial effusions occurring in patients who underwent randomization within 1 week following surgery. Therefore, a delayed NOAC initiation strategy may have mitigated the risk of major bleeding. Despite these unfavorable results, ongoing research on NOACs as a lone anticoagulation strategy is currently underway with a phase II study examining the use of Rivaroxaban in patients
with mechanical prostheses (CATHAR, NCT02128841). Additionally, a non-inferiority trial comparing a low INR strategy (1.5–2) and NOACs in patients with previous AVR using the On-X prosthesis should be considered.

In conclusion, the PROACT trial demonstrates the safety and feasibility of targeting lower INR values in patients with home anticoagulation monitoring undergoing mechanical AVR using the On-X prosthesis. This represents a significant step forward in patient management, with demonstrable improvements in patient outcomes. Findings from this trial highlight the need for a prospective randomized trial comparing mechanical and bioprosthetic valves to the Ross procedure in young adults undergoing AVR.

COMPETING INTERESTS

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