Randomized Clinical Trial
Scoliocorrector Fatma-UI For Correction of Adolescent Idiopathic Scoliosis: Development, Effectivity, Safety and Functional Outcome

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Abstract
BACKGROUND
Adolescent idiopathic scoliosis remains a major problem due to its high incidence, high risk, and high cost. One of the aims of the management in scoliosis is to correct the deformity. Many techniques are available to correct deformity in scoliosis. However, all of them are far from ideal to achieve three-dimensional correction in scoliosis.

AIM
The aim of this study is therefore to develop a set of tools named Scoliocorrector Fatma-UI (SCFUI) to aid three-dimensional correction and to evaluate the efficacy, safety, and functional outcome.

METHODS
This study consists of two stages. In the first stage, we developed the SCFUI and tested it in finite element and biomechanical test. The second stage was a single-blinded randomized clinical trial to evaluate the SCFUI compared to direct vertebral rotation (DVR). Forty-four subjects with adolescent idiopathic scoliosis were randomly
allocated into DVR group ($n = 23$) and SCFUI group ($n = 21$). Radiological, neurological, and functional outcome was compared between group.

RESULTS
Finite element revealed maximum stress of the SCFUI components are between 31.2-252 MPa. Biomechanical analysis revealed the modulus elasticity of SCFUI was 95 613.24±6 332.77 MPa. Both groups show improvement in Cobb angle and sagittal profile, however the RaSag angle was lower in SCFUI group (11.59±7.46 vs 18.23±6.39, $P = 0.001$). Neurological and functional outcome were comparable in both groups.

CONCLUSION
We concluded that SCFUI developed in this study resulted in similar coronal and sagittal but better rotational correction compared to DVR. The safety and functional outcomes were also similar to DVR.

INTRODUCTION
Adolescent idiopathic scoliosis (AIS) remains a major problem due to its high incidence, high risk, and high cost. The incidence of AIS reaches as high as 13%.$^1$ Management of AIS is also subject to a high risk of complications such as death, pseudoarthrosis, infection, neurological deficits, and pedicle screw misplacements.$^2$ Cost needed to manage AIS reached USD 55,000 in 1997 and USD 177,000 in 2012 and keep increasing.$^3$ One of the aims of management in adolescent idiopathic scoliosis is to correct the deformity. While coronal correction remains the major aim, corrections in sagittal and axial planes are also necessary. Both sagittal and axial plane deformity are associated with the decrease in lung function.$^{4,5}$
Many techniques are available to correct deformity in scoliosis. However, all of them are far from ideal to achieve three-dimensional correction in scoliosis.$^6$ Direct vertebral rotation gives excellent coronal and rotational correction but causes thoracic
hypokyphosis. Its safety is also questioned. Posteromedial translation is reported to result in good coronal and sagittal correction but its rotational correction is questioned. Sublaminar fixation point in posteromedial translation is located posterior to the rotational axis in scoliosis and may affect the axial correction. Posteromedial translation also requires a universal clamp and sublaminar band which are not available in many countries.

The aim of this study is therefore to develop a set of tools named Scoliocorrector Fatma-UI (SCFUI) to aid three-dimensional scoliosis correction and to evaluate the efficacy, safety, and functional outcome of the tools in AIS surgery.

MATERIALS AND METHODS

This study consists of two stages. The first stage was the development of the SCFUI, as well as finite element and biomechanical analysis to evaluate the validity and safety of the SCFUI. The second stage was a single-blinded randomized clinical trial to evaluate the effectivity, safety, and functional outcome of the SCFUI compared to direct vertebral rotation.

Stage I. SCFUI development

Development of SCFUI started in January to May 2020 in the Integrated Creative Learning Laboratory, Faculty of Technics, University of Indonesia. We listed anticipated benefits of the tools as well as ways to achieve them and formulated them in conceptual design. The conceptual design was subsequently developed into a technical design with an initial size complying with pedicle screws available in the market. Technical design was developed in Solidworks v.2017 software. The design was tested in finite element analysis using ANSYS v.2020 software. We tested the design by static structural method, no separation contact, automatic mesh, and stainless-steel 316L material with a testing force of 800N. We modified the size and design until the stress was lower than the yield strength of stainless steel 316L. Using Ultimaker Cura v.11, we modified the size and orientation of the SCFUI design to comply with the building machine. SCFUI was built using CNC milling Harford 1000x500x500 dan CNC Milling Eccoca
The prototype was then tested using Universal Testing Machine Tensilon RTF 2350. Four hundred Newton's force was used to pull SCFUI with a speed of 50 mm/s for ten times and the modulus of elasticity was recorded. Stress-strain curve was also created. We also pull the prototype until the failure point to observe the maximum force as well as the response at the failure point.

Stage 2. Clinical trials

The second stage of the study was conducted in Fatmawati General Hospital from June 2020 to June 2021. Inclusion criteria were patients with adolescents aged 11 to 18 with Lenke 1 idiopathic scoliosis 40 - 100 degrees, without a history of previous spinal surgery. Forty-four- subjects were randomized to the intervention group (SCFUI, n = 21) and control group (DVR, n = 23) using Randlist® software. The subject was blinded to the randomization. Ethical clearances from the Faculty of Medicine University of Indonesia number 615/UN2.F1/ETIK/PPM.00.02/2020 and Fatmawati General Hospital number DM01.01/VIII.2/294/2020 were obtained before the study.

Under general anesthetic and intraoperative neuromonitoring, the subject was put prone on two bolsters in the chest and pelvis region and fixed to the surgical table. The spine was exposed from the upper to lower instrumented vertebra (LIV). The upper instrumented vertebra (UIV) was the neutral vertebra cranial to upper-end vertebra while LIV is the last significantly touched vertebra. Screws were inserted in UIV, 1 Level distal to UIV, apical vertebra, 1 Level proximal and 1 Level distal to apical vertebra, 1 Level proximal to LIV and LIV. Whenever there were more than three subsequent vertebrae to UIV or LIV, additional screws were put until there were no more than three instrumented subsequent vertebrae. The screws were inserted free hand, without any specific attempt to hit the cortical bone in front of the vertical body as it will increase risk of injury. A correction was performed according to the group allocation, and the wound was closed layer by layer.

For the intervention group, five pulleys were attached to the apical, one level above and below the apical screw, and to the closest screw proximal and distal to the apical screws. The correction board was attached to the surgical table on the concave side. Five
correction screws and their housing were assembled and attached to the correction board according to the normal sagittal profile and intended direction of pulling. The lowest screw housing should be at least 5 cm higher than the most prominent point of the curve. Five wires were fixed to the correction screws and housings and circled to the pulleys to form moveable pulleys. A correction was achieved by gradually and alternately rotating the correction screws to pull the moveable pulleys. The pull was stopped when correction was achieved, screw rotation was maximized, or whenever there was a screw pullout. The Convex rod was attached to secured with set screws. Pulleys were removed and the concave rod was attached and secured with set screws.

For the control group, both rods were inserted and loosely secured with set screws. Tubes were inserted into the screws closest to the apical vertebrae. The vertebrae were rotated as neutral as possible and set screws were tightened. Other tubes were inserted into apical dan periapical screws and rotated while counter-rotation was performed on the distal neutral tubes. All set screws were tightened.

For radiological outcome we measured coronal cobb angle and T5 - 12 kyphosis angle from a standing radiograph, as well as RaSag of the apical vertebra from CT - Scan. The radiograph and CT - Scan were obtained preoperatively and within 10 days after surgery. Intraoperative MEP, SSEP, and EMG as well as post-operative motor power of the lower extremity were also recorded. Functional outcome was also measured using the SRS-22 questionnaire. The data were compared between groups and statistical analysis was performed using STATA v. 14.

**RESULTS**

Stage I. SCFUI Development

We first identified the needs and then the principles and methods to fulfill the needs. The concept is shown in Table 1. Based on the concept we developed a set of tools consisting of a correction board, correction screw, and its housing, wire, and pulley. The
pulley would be attached to pedicle screws and pulled toward correction screws secured by screw housing on the correction board. (Figure 1) From the finite element analysis, the maximum stress of the correction board, correction screw, screw housing, and wire were 126.5 MPa, 31.2 MPa, 92.1 MPa, and 252 MPa respectively. (Figure 2) Passing finite element analysis, the SCFUI prototype was built. (Figure 3) Biomechanical analysis revealed the modulus elasticity of SCFUI was 95 613,24±6 332,77 MPa. The stress-strain curve is shown in Figure 4. Failure of SCFUI was observed on 800 Nm as detachment of wire from correction screw.

Stage 2. Clinical study
Baseline characteristics of research subjects are shown in Table 2 while corrections using intervention and control groups are in Figure 5. Body weight, coronal curve, and RaSag in the control group were higher than in the intervention group.

Postoperatively, only RaSag in the control group was higher than the intervention group. (Table 4.3 and Figure 6) In both groups, no abnormality of motor-evoked potential, somatosensory-evoked potential, and neurotonic discharge was found during surgery. (Figure 7) Postoperatively, no motor deficit was found in all groups. During the study, no adverse event, screw pullout, or wire breakage was observed.

DISCUSSION
In the development of SCFUI, stainless-steel 316L was used as the material. Stainless-steel 316L is strong, stain-resistant, low maintenance, and economical.\textsuperscript{10} It is also one of the most used alloys for spinal implants.\textsuperscript{11,12}

In finite element analysis, the highest stress was observed in the wire. However, the stress was lower than the yield strength and ultimate strength of stainless steel 316L.\textsuperscript{13} It indicates that SCFUI is in the elastic phase, meaning that it can withstand force without any change in shape or failure.\textsuperscript{10,13} During the analysis, the force used is far higher than the force required in clinical application or the force to cause the pullout of the screw.\textsuperscript{13} Moveable pulley configuration in SCFUI also gives a mechanical advantage to lower the force needed to correct the deformity up to 200%.\textsuperscript{14} Biomechanical studies validated the
finding of finite element analysis. The modulus elasticity of SCFUI is lower than the modulus elasticity of stainless-steel 316L and the stress-strain curve is proportional. Hooke law describes if the stress is proportional to the strain, a material is in elastic condition, thus it will return to its initial shape whenever the force is removed.

At the point of failure, the wire attachment to the correction screw will be detached slowly. The detachment occurs slowly and can be observed with the naked eye. The finding emphasizes the safety of SCFUI in the event in the point of failure, it will not endanger the patient.

The SCFUI is therefore compliant with the ASTM F2193 standard. ASTM F2193 is the international standard for the development of surgical instruments for spinal surgery. It included the design, strength, and safety of the instrument.

Coronal correction is important in the evaluation of surgery outcomes. For patients, coronal correction is the main indicator of the success of surgery. Medically, coronal correction is correlated to functional outcome. In SCFUI, pedicle screw fixation is the use of higher force for correction. Moreover, the moveable pulley increases the force acting on the spine to two folds. It also optimized the vector of pull.

Our coronal correction is similar to correction using posteromedial translation by Mazda et al. They reported coronal correction of 66±13%. A meta-analysis reported the coronal correction of scoliosis surgery was 60-80% despite techniques used for correction.

Evaluation of the sagittal profile is also important. An imbalance of the sagittal profile will result in poor pulmonary function and increased complications. In our study, both techniques gave a good sagittal profile. The reported hypokyphosis due to DVR was not observed in our study.

SCFUI uses the correction board as the pulling point for translation. It allows a longer distance of posteromedial translation. However, the advantage of SCFUI over DVR for sagittal correction was not proven in our study. The advantage of SFUI for sagittal correction is negated by the use of pedicle screws which also cause hypokyphosis. Moreover, the DVR in our study also resulted in good sagittal correction.
Rotational correction nowadays is also emphasized in scoliosis correction. Poor rotation will result in rib cage asymmetry and decreased respiratory function. In our study, a better rotation correction was achieved by SCFUI. SCFUI utilized pedicle screw fixation that penetrate the vertebral body, thus allowing a fixation axis anterior to the rotation axis in scoliosis. Fixation points anterior to the rotational axis in mandatory to achieve adequate torsion for vertebra derotation. However, it should be noted that our result is subject to bias as a bigger rotation was found in the control group before the study.

We did not find any neurological deficits in our study. It might indicate that both techniques were safe. Nevertheless, scoliosis in our study was limited to a mild curve in which the incidence of neurological deficit is quite low. The incidence of the neurological deficit will be increased by the severity of the curve and the presence of hyperkyphosis. None of the subjects in our study had hyperkyphosis. SCFUI was developed with a novel design and complied with the international standard for spinal instruments. The efficacy, safety, and functional outcomes had also been evaluated in the clinical study. However, our study was subject to several limitations. We did not measure the duration needed to assemble the correction tools until the correction was performed, nor the bleeding occurred during the period. SCFUI is quite heavy and the assembling takes time. Further study is necessary to simplify the assembling and to reduce the size as well as the weight. Scoliosis evaluated in this study was also very narrow, limited to Lenke type 1 curve only. Further study involving wider Lenke criteria should be performed.

**CONCLUSION**

We concluded that SCFUI developed in this study comply with the international standard for spinal instrument. It resulted in similar coronal and sagittal but better rotational correction compared to DVR. The safety and functional outcomes were also similar to DVR.
ARTICLE HIGHLIGHTS

Research background
Adolescent idiopathic scoliosis remains a main problem in orthopedic due to its high incidence, high risk, and high cost.

Research motivation
Ideal correction of scoliosis should be three-dimensional. However, many techniques failed to achieve three-dimensional correction.

Research objectives
We developed a novel set of tools to aid three-dimensional correction of adolescent idiopathic scoliosis.

Research methods
We performed finite element and biomechanical test for the tools, and evaluated its radiological, neurological and functional outcomes in a randomized clinical trial to ensure its efficacy and safety.

Research results
The tools developed passed the finite element analysis and biomechanical test as well as clinical study.

Research conclusions
We developed a novel set of tools to aid three-dimensional correction of adolescent idiopathic scoliosis.

Research perspectives
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