# Circum-Psoas Block versus Supra-Inguinal Fascia Iliaca Block for Postoperative Analgesia in Patients Undergoing Total Hip Arthroplasty: A Randomized Clinical Trial

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**Purpose:** Total hip arthroplasty (THA) is often associated with moderate to severe pain. The present study compared the efficacy of circum-psoas block (CPB) with supra-inguinal fascia iliaca block (SIFIB) for postoperative analgesia in patients undergoing THA. **Patients and Methods:** In this randomized trial, sixty-four patients undergoing THA were allocated randomly to the CPB group or SIFIB group with 40 mL of 0.3% ropivacaine. The primary outcome was dynamic pain score at 6 h postoperatively. Secondary outcomes included dynamic pain scores at 12, 24 and 48 h; static pain scores; sensory and motor block; opioid consumption; time to first opioid request; length of hospital stay; patient satisfaction; and adverse events.

**Results:** CPB patients showed significantly lower dynamic pain scores at 6 (3.11  $\pm$  0.66 vs 4.47  $\pm$  0.74, respectively; P = 0.000), 12 (2.52  $\pm$  0.73 vs 3.53  $\pm$  0.85, respectively; P = 0.000) and 24 h (2.30  $\pm$  0.57 vs 2.87  $\pm$  0.71, respectively; P = 0.001) after surgery, as well as lower static pain scores at 6 and 12h (P = 0.001 and P = 0.033 respectively) than SIFIB patients. Lower opioid consumption was observed in the CPB group at 24 and 48 h (P = 0.000, both) than in the SIFIB group. Patients in the CPB group reported improved quadriceps strength at 6 and 12 h (P = 0.000, both), as well as better muscle strength of hip flexion at 6, 12 and 24 h (P = 0.000, P = 0.000 and P = 0.025 respectively). Compared with SIFIB, CPB was associated with increased sensory block coverage at 6, 12 and 24 h (P = 0.000, P = 0.000, and P = 0.022, respectively).

Conclusion: CPB has a greater potential to alleviate postoperative pain and improve recovery in THA patients than SIFIB.

Keywords: total hip arthroplasty, lumbar plexus, sacral plexus, regional anesthesia

#### Introduction

Total hip arthroplasty (THA) is a common surgical procedure, with more than 280,000 THA procedures performed annually in the United States. Patients undergoing THA often experience moderate to severe pain, which can reduce their overall quality of life and even develop into persistent pain. Consequently, efforts are made to facilitate enhanced recovery profiles to reduce postoperative pain in THA patients. At present, multimodal analgesia approaches have been used for pain management after THA, including oral analgesia, epidural analgesia, and peripheral nerve block. However, pain management after THA is a challenging goal to achieve, and an ideal analgesic regimen for THA has yet to be defined.

Hebbard et al first proposed ultrasound-guided supra-inguinal fascia iliaca block (SIFIB) in 2011.<sup>7</sup> As part of a multimodal analgesic approach, it is now frequently used for THA.<sup>8,9</sup> With the cranial spread of local anesthetic (LA), SIFIB has been suggested to effectively block the femoral nerve (FN), the obturator nerve (ON) and the lateral femoral cutaneous nerve (LFCN).<sup>10</sup> However, the SIFIB approach often leads to quadricep weakness<sup>11,12</sup> and fails to block the subcostal nerve, iliohypogastric nerve, ilioinguinal nerve, and sacral nerves, which are considered to be highly

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important for postoperative analgesia after THA. <sup>13–15</sup> In addition, Bendtsen et al argued that the ON did not cross the fascia iliaca compartment, <sup>16</sup> thus compromising the effectiveness of the obturator nerve block caused by SIFIB.

Recently, Li et al reported a new circum-psoas block (CPB) technique.<sup>17</sup> With this approach, sensory block coverage can be achieved from the T8-11 to S1-3 dermatomes with little effect on quadriceps strength. This may help to provide more widespread analgesia coverage than the SIFIB approach in patients with THA.

We thus conducted this study to assess whether CPB is associated with superior pain control and reduced opioid consumption compared to SIFIB among patients undergoing THA. The primary outcome was dynamic pain score at 6 h after surgery.

### **Materials and Methods**

The current trial was approved by the Ethics Committee Board of Beijing Chaoyang Hospital, Capital Medical University on October 11, 2021, with reference number 2021-ke-583 and registered at <a href="http://www.chictr.org.cn">http://www.chictr.org.cn</a> (ChiCTR2100053421) on November 20, 2021. The study adhered to the Consolidated Standards of Reporting Trials (CONSORT) guidelines and Helsinki declaration.

### **Participants**

We enrolled patients between December 31, 2021, and September 10, 2022, who would undergo elective THA with posterolateral approach under spinal anesthesia (Figure 1). The study protocol was explained to eligible patients, and they

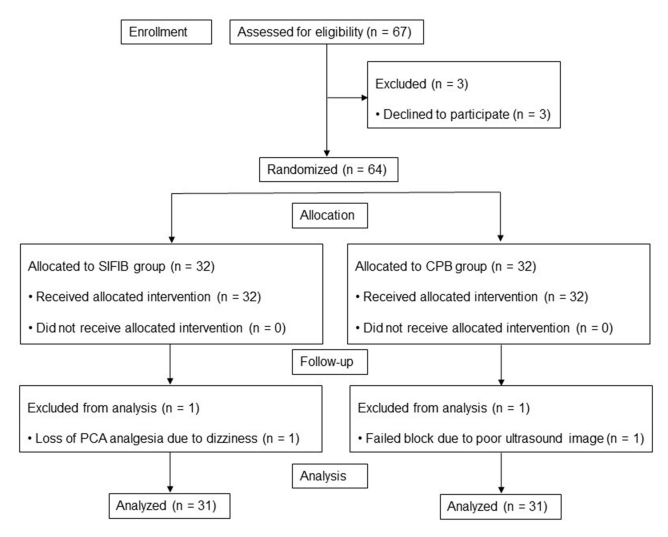


Figure I The CONSORT study flowchart.

Abbreviations: CONSORT, Consolidated Standards of Reporting Trials; PCA, patient-controlled analgesia.

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provided written informed consent to our team. The study included participants aged 18-90 with a body mass index (BMI) of 18-30 kg/m<sup>2</sup> and American Society of Anesthesiologists (ASA) physical status class I to III. Patients with puncture site infection, anticoagulant status, allergy to LA, serious cardiovascular disease and blood disease, chronic opioid consumption or alcohol abuse, myasthenia gravis or Parkinson's disease or pregnancy, previous hip or spinal surgery history, narrow interspace between the transverse process of L5 and the sacral ala (X-ray film anteroposterior: less than 1 cm), cognitive dysfunction before surgery and inability to cooperate with pain assessment were excluded.

### Randomization and Blinding

Sixty-four eligible participants were randomly assigned to receive either SIFIB or CPB by computerized sequence generation. Randomization and group allocation were performed at a 1:1 intergroup ratio, and allocation information was placed in opaque sealed envelope. Two investigators blinded to the study design opened the sealed envelope just before surgery and prepared a syringe containing 40 mL of 0.3% ropivacaine that was labeled with the randomization number of the patient receiving the block. Afterward, a skilled anesthesiologist performed the ultrasound-guided nerve block according to the allocation. All other researchers and outcome assessors were masked to the allocation of the groups until the study was completed.

### Intraoperative Management

In the preoperative holding area, all patients received celecoxib 200 mg and acetaminophen 650 mg orally. After entering the operating room, all patients were monitored with standard ASA monitors (including pulse oximetry, noninvasive blood pressure, and electrocardiography), and peripheral venous access was obtained. Then, an arterial catheter was inserted into the radial artery for haemodynamic monitoring.

After premedication with 1–2 mg midazolam, patients were placed in a lateral decubitus position with the operative side up to receive spinal anesthesia. Hypobaric ropivacaine (2.5 mL of 0.5% ropivacaine) was administered intrathecally at the L3-4 interspace, which was injected over 30s. Patients remained for 15 minutes in a lateral decubitus position to achieve a more intense block of the operative side. Spinal anesthesia was considered to be successful if sensory block to T10 level was acquired, <sup>18</sup> as evaluated by loss of cold sensation. For both blocks, a 100-mm, 22-gauge needle (Tuoren, Henan, China) was inserted with strict antiseptic protocols. All procedures were performed under ultrasonographic guidance (KONICA Minolta, Tokyo, Japan) by trained regional anesthesia specialists (YW, JZ, and HL), who performed CPB and SIFIB in more than 25 patients in the preliminary study.

For SIFIB, patients were placed supine after 15 minutes of spinal anesthesia. A linear transducer was used to obtain the image of the anterior superior iliac spine in the sagittal position. The internal oblique muscles, iliacus and sartorius can be identified by sliding the probe medially (Figure 2). After identifying the "bow-tie sign", the needle was inserted in a caudad-cephalad direction. Using an in-plane approach, the fascia iliaca was penetrated and hydrodissected from the iliacus muscle with 0.9% saline (2-5 mL). Upward movement of the deep circumflex artery after injection was a sign of a successful injection beneath the fascia iliaca. Following repeated negative aspiration, 40 mL of 0.3% ropivacaine was injected into the fascia iliaca compartment.

For CPB, the procedure was performed with lateral decubitus after 15 minutes of spinal anesthesia. A curved array transducer was placed in a paramedian sagittal plane to identify the sacrum and the transverse process of the L5 vertebral body of the surgical side. In this position, the probe was rotated 90 degrees to obtain the paramedian transverse plane of the lumbosacral region until the intertransverse ligament, and the lumbosacral ligament could be seen in the sonogram between the iliac bone and laminae of L5 (Figure 3A). Using an out-of-plane approach, the needle was inserted to penetrate the erector spinae, the intertransverse ligament, and the lumbosacral ligament. An endpoint of injection was an observation of the needle tip penetrating the lumbosacral ligament and a distinct loss of resistance. <sup>19</sup> Injection of 2–5 mL of 0.9% saline was used to confirm the correct position of the needle tip. An additional 20 mL of 0.3% ropivacaine was injected with repeated negative aspiration. After that, the curved array transducer was placed cranial to the iliac crest in a transverse plane and slid to identify the shamrock sign. A puncture needle was then inserted in a posterolateral to anteromedial direction with an in-plane approach until the needle tip penetrated the transversalis fascia covering the psoas major (PM) muscle, and a loss of resistance was obtained (Figure 3B). The distribution of fluid around the antero-lateral margin of the PM muscle beneath the transversalis fascia was used as an

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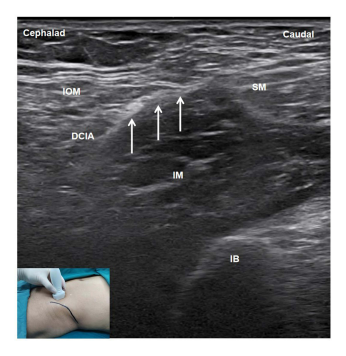


Figure 2 Sonoanatomy of the supra-inguinal fascia iliaca block.

Note: The white arrows indicate fascia iliaca.

Abbreviations: IOM, internal oblique muscle; DCIA, deep circumflex iliac artery; SM, sartorius muscle; IM, iliac muscle; IB, iliac bone.

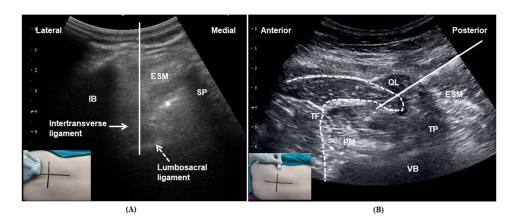


Figure 3 (A) Sonoanatomy of the circum-psoas block at the L5/S1 intervertebral level. (B) Sonoanatomy of the circum-psoas block at the supra-iliac level. Notes: (A) The white solid line indicates the needle trajectory. The solid arrow indicates the intertransverse ligament. The dashed arrow indicates the lumbosacral ligament. The asterisk indicates the laminae of L5. (B) The white solid line indicates the needle trajectory. The white dashed line indicates the TF. Abbreviations: IB, iliac bone; ESM, erector spinae muscle; SP, spinal process; TF, transversalis fascia; QL, quadratus lumborum; PM, psoas major; VB, vertebral body; TP, transverse process.

indicator of successful penetration of the transversalis fascia following injection of 2-5 mL of 0.9% saline. With intermittent negative aspiration, an additional 20 mL of 0.3% ropivacaine was injected. In this study, all surgical interventions were performed by the same team of orthopaedic surgeons from our institution, and the duration of surgery was recorded.

# Postoperative Pain Management

After surgery, patients were provided with a patient-controlled analgesia (PCA) pump, which was programmed to deliver 2 µg of intravenous sufentanil on demand with a lockout interval of 15 min. In the surgical ward, in addition to PCA, they also received regular doses of celecoxib 200 mg orally every 12 h and acetaminophen 650 mg orally every 8

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h. Rescue analgesics (administered if patients could not tolerate the pain) consisted of oxycodone 5 mg administered orally in the ward.

#### Assessment of Outcomes

The primary outcome was dynamic (with passive 90° flexion of the hip) pain score at 6 h after surgery. Pain intensity was evaluated using a visual analogue scale (VAS) ranging from 0 (no pain) to 10 (extreme pain).

Secondary outcomes included opioid consumption at 24 and 48 h (expressed as intravenous morphine milligram equivalents (MME) using a conversion factor of 1000<sup>20</sup> and 1.5<sup>21</sup> for intravenous sufentanil and oral oxycodone, respectively); dynamic pain scores at 12, 24 and 48 h; static (at rest) pain scores, sensory block, motor block at 6, 12, 24 and 48 h; first opioid request; length of hospital stay (the time from randomization to hospital discharge); patient satisfaction (0 = "unsatisfied", 10 = "extremely satisfied"); and incidence of adverse events (including nausea, vomiting, dizziness, urinary retention, respiratory depression, vascular puncture, nerve injury, and local anesthetic toxicity).

Sensory block was evaluated by using the standardized stimuli of ice and compared with the contralateral side. Rating was categorized as normal sensation and any changes in sensation (eg, no sensation or decreased sensation). The test was conducted at the anterior axillary line to determine the blocked areas of thoracic spinal nerves.<sup>22</sup> Lumbosacral nerves (L1-S3), FN and LFCN were measured at different dermatomes according to a previous study.<sup>23</sup>

Motor block was evaluated by measuring the muscle strength of hip flexion, knee extension, and ankle flexion. The following criteria were used to score muscle strength: 0, no muscle contraction; 1, muscle contraction without joint movement; 2, joint movement but no gravity resistance; 3, movement against gravity; 4, gravity and partial against resistance; and 5, normal strength.

### Statistical Analysis

Based on a previous study, the median minimal clinically important difference (median MCID) for dynamic pain score was absolute 1.8 after THA.<sup>24</sup> This study was powered to detect a difference of 2 in dynamic pain scores<sup>21</sup> between patients undergoing CPB versus those receiving SIFIB, assuming a SD of 2.5. Using a two-sample t test, a sample of 52 participants (26 in each group) would have approximately 80% power to detect a clinically significant reduction in pain score with a significance level alpha of 0.05. Taking into account a dropout rate of approximately 20%, a total of 64 patients were selected with 32 patients per group.

Statistical analysis was performed using the SPSS V.19 software package (IBM, Armonk, New York). For continuous data, normality was first assessed with the Shapiro-Wilk normality test. Data with a normal distribution were reported as mean [standard deviation (SD)] and analyzed with Student's t test, while nonnormal data and ordinal data were expressed as median [interquartile range (IQR)] and analyzed with the Mann-Whitney U-test. Categorical data were compared with the chi-squared  $(\chi^2)$  test. When the expected count for any of the aforementioned categorical data was less than five, Fisher's exact test was used. A two-sided P value of <0.05 was considered as statistically significant.

#### Results

The flowchart according to the CONSORT statement is shown in Figure 1. In the present study, 67 patients were enrolled, and three were excluded due to refusal to participate. Consequently, we randomly assigned 64 patients to either the CPB or SIFIB groups, each with 32 patients. One SIFIB group patient withdrew from PCA analgesia due to dizziness. The block failed in one CPB group patient due to poor ultrasound image quality. Finally, 62 patients completed the study and were analyzed (n = 31 in each group). The demographic parameters, surgical side and surgical duration are presented in Table 1.

# **Primary Outcome**

As shown in Table 2, patients receiving CPB had lower dynamic pain scores than those receiving SIFIB at 6 hours after surgery (3.11  $\pm$  0.66 vs 4.47  $\pm$  0.74, respectively), reaching statistical significance (P = 0.000).

Table I Demographic and Clinical Characteristics

Variable	SIFIB (n = 31)	CPB (n = 31)	P-value
Age (years)	68.16 ± 5.91	68.90 ± 6.12	0.629
Gender (M/F)	14 (45.2%)/17 (54.8%)	16 (51.6%)/15 (48.4%)	0.611
Height (cm)	163.97 ± 6.96	165.97 ± 6.52	0.248
Weight (kg)	65.71 ± 8.89	65.52 ± 9.80	0.935
BMI (kg/m <sup>2</sup> )	24.39 ± 2.89	23.70 ± 2.65	0.328
ASA (I–II/ III)	20 (64.5%)/11 (35.5%)	22 (71.0%)/9 (29.0%)	0.587
Surgery side (Left/ Right)	15 (48.4%)/16 (51.6%)	18 (58.1%)/13 (41.9%)	0.445
Surgical duration (min)	85.19 ± 14.05	83.77 ± 14.37	0.696

Note: Data are mean ± SD, or number of patients (%)

Abbreviations: ASA, American Society of Anesthesiologists; BMI, body mass index; CPB, circum-psoas block; SIFIB, supra-inguinal fascia iliaca block.

Table 2 Postoperative Pain Scores, Opioid Consumption, and Number of Affected **Dermatomes** 

Variable	CPB (n = 31)	SIFIB (n = 31)	P-value
Pain scores			
Pain at 6 h (dynamic)	3.11 ± 0.66	4.47 ± 0.74	0.000
Pain at 12 h (dynamic)	2.52 ± 0.73	3.53 ± 0.85	0.000
Pain at 24 h (dynamic)	2.30 ± 0.57	2.87 ± 0.71	0.001
Pain at 48 h (dynamic)	2.10 ± 0.59	2.37 ± 0.61	0.077
Pain at 6 h (rest)	1.42 ± 0.61	2.05 ± 0.83	0.001
Pain at 12 h (rest)	1.27 ± 0.47	1.61 ± 0.72	0.033
Pain at 24 h (rest)	0.98 ± 0.40	1.19 ± 0.50	0.073
Pain at 48 h (rest)	0.77 ± 0.40	0.87 ± 0.42	0.359
Time to first opioid request (h)	7.0 (5.5–9.5)	5.5 (4.0–9.5)	0.048
Opioid consumption at 24 h (mg)	6.0 (4.0-8.0)	9.3 (8.0–12.0)	0.000
Opioid consumption at 48 h (mg)	7.33 (4.0–10.0)	11.33 (9.3–15.3)	0.000
Number of affected dermatomes			
At 6 h	9 (9–10)	4 (4–4)	0.000
At I2 h	7 (5–9)	3 (3-4)	0.000
At 24 h	2 (0–6)	I (0-2)	0.022
At 48 h	0 (0–0)	0 (0–0)	0.665

Notes: Continuous variables are presented as mean ± SD; nonnormal data are presented as median (IQR). Abbreviations: CPB, circum-psoas block; SIFIB, supra-inguinal fascia iliaca block.

# Secondary Outcomes

In the CPB group, dynamic pain scores were significantly lower at 12 h and 24 h (P = 0.000 and P = 0.001 respectively), as well as static pain scores at 6 h and 12 h (P = 0.001 and P = 0.033 respectively) after surgery (Table 2). The VAS scores of the two groups were not significantly different at 48 h after surgery. Besides, the cumulative opioid requirements were lower in the CPB group at 24 h and 48 h (P = 0.000, both) than in the SIFIB group (Table 2). There were no differences in the proportion of patients who needed oxycodone at 24 (CPB 3.2% vs SIFIB 16.1%, P = 0.2) and 48 h (CPB 3.2% vs SIFIB 19.4%, P = 0.1).

Compared with SIFIB, CPB was associated with reduced motor block of the quadriceps at 6 and 12 h (P = 0.000, both), as evidenced by enhanced knee extension (Table 3). Furthermore, CPB significantly improved the muscle strength of hip flexion at 6, 12 and 24 h (P = 0.000, P = 0.000 and P = 0.025 respectively). Patients in the CPB group reported median (IQR) muscle strength with ankle flexion at 6 h of 5 [5 to 5] compared to 5 [5 to 5] in the SIFIB group (P = 0.088). The muscle strength of ankle flexion was not reported at 12, 24 and 48 h after surgery since it completely regressed at 12 h in all patients.

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Table 3 Postoperative Motor Block, Adverse Events, Length of Hospital Stay, and Satisfaction Score

Variable	<b>CPB</b> (n = 31)	SIFIB (n = 31)	P-value
Motor block			
Knee extension at 6 h	4 (3–4)	3 (2–4)	0.000
Knee extension at 12 h	4 (4–5)	3 (3–4)	0.000
Knee extension at 24 h	5 (5–5)	5 (4–5)	0.375
Knee extension at 48 h	5 (5–5)	5 (5–5)	0.691
Hip flexion at 6 h	4 (4-4)	3 (2–3)	0.000
Hip flexion at 12 h	4 (4–5)	3 (3–4)	0.000
Hip flexion at 24 h	5 (4–5)	4 (4–5)	0.025
Hip flexion at 48 h	5 (5–5)	5 (4–5)	0.122
Ankle flexion at 6 h	5 (5–5)	5 (5–5)	0.088
Nausea [n (%)]	2 (6.5%)	3 (9.7%)	1.000
Vomiting [n (%)]	I (3.2%)	I (3.2%)	1.000
Dizziness [n (%)]	0 (0)	I (3.2%)	1.000
Urinary retention [n (%)]	I (3.2%)	2 (6.5%)	1.000
Length of hospital stay (d)	6 (5–6)	6 (5–7)	0.476
Patient satisfaction score	10 (9–10)	10 (9–10)	0.810

Note: Data are presented as median (IQR) or number of patients (%). Abbreviations: CPB, circum-psoas block; SIFIB, supra-inguinal fascia iliaca block.

The presence of sensory blockade for different dermatomes and for the terminal nerves are shown in Figure 4. Compared with SIFIB, CPB was associated with significantly increased dermatomal coverage of sensory block at 6, 12 and 24 h (P = 0.000, P = 0.000 and P = 0.022 respectively).

As shown in Table 3, patient satisfaction scores were high between groups with no significant difference. There was no difference in length of hospital stay (median, IQR) - CPB group 6 [5 to 6] days, SIFIB 6 [5 to 7] days. No differences

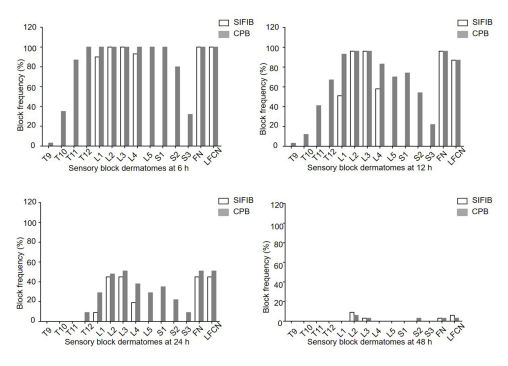


Figure 4 The sensory block probability of different dermatomes and terminal nerves at different time point between groups. Abbreviations: SIFIB, supra-inguinal fascia iliaca block; CPB, circum-psoas block.

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were found in the incidence of nausea, vomiting, dizziness, or urinary retention after surgery between the two groups. No patients in either group suffered clinically significant vascular injury, nerve injury, respiratory depression, or local anesthetic toxicity.

### **Discussion**

This prospective randomized trial compared ultrasound-guided CPB with SIFIB in patients undergoing THA. We found that patients in the CPB group had significantly lower dynamic pain scores at 6 h with improved quadriceps strength when compared with SIFIB. These findings support the use of CPB as an effective approach to promote rehabilitation in THA patients.

Although Diwan et al initially described circumpsoas block<sup>25</sup> by depositing LA between the anterior psoas fascia and the PM muscle, this method can only block nerves emerging from the lumbar plexus. Due to this, Li et al proposed another method for CPB in the previous study, 17 which differed from Diwan's method in terms of injection sites and dermatomal coverage. In the present study, we thus observed extensive sensory block coverage ranging from the T9-12 to S1-3 dermatomes after CPB.

Our primary outcome requires discussion, as a number of clinical studies 15,26,27 have chosen opioid consumption as pain surrogate marker since the use of PCA devices became common practice. However, early mobilization with low level of pain intensity is essential for enhanced postoperative recovery. In our study, CPB was associated with significantly lower dynamic pain score compared with SIFIB (3.11  $\pm$  0.66 vs 4.47  $\pm$  0.74) at 6 h postoperatively. Although we did not detect a difference of 2 in dynamic pain score, a 31% decrease in dynamic pain score still echoes the median MCID for a relative pain reduction recommended by Laigaard et al. 24 This encouraging result can be attributed to the extensive sensory block coverage of the CPB approach.

We additionally observed lower dynamic pain scores at 12 and 24 h, lower static pain scores, and less opioid consumption postoperatively in patients who received CPB compared with those who received SIFIB. However, the differences between CPB group and SIFIB group may be of limited clinical importance, as they failed to meet the median MCIDs for pain scores and opioid consumption.<sup>24</sup> Despite all this, CPB can provide practitioners with alternative analgesic options for managing postoperative pain in patients undergoing THA. In our study, we found no differences in rescue analgesia need between the groups. The result may partly be due to the limited sample size. Furthermore, PCA pumps were used to provide immediate pain relief when pain gradually increased, thus reducing the number of patients requiring rescue analgesia in the SIFIB group.

We also found that CPB led to sensory blockage but had little effect on motor function. This observation may be due to the fact that we use relatively low concentration of ropivacaine. In addition, the nerves are located away from the inject site of CPB block, so ropivacaine hydrochloride could be dissipated during the spread of the LA near the target nerves, which may result in a lower concentration of LA acting on nerves than we actually used. Therefore, CPB is effective in improving the early recovery of motor function by preserving muscle strength. Notably, however, the muscle strength of ankle flexion did not differ between groups. Since CPB results in sensory block at level L5/S1, one should expect an inherent impairment of ankle mobilization as well. The lack of intergroup differences may be explained by the fact that ropivacaine has the property of separation between motor and sensory blockade at low concentrations; thus, our trial may have not been able to fully evaluate the changes in muscle strength using the current method.

CPB appears to be safe in the present study, as no block-related complications were observed. Despite being in a deep location, CPB is situated away from major blood vessels, which reduces the risk of complications. In addition, sonography can be used to identify the needle position and prevent vascular puncture. In our trial, CPB was performed with a relatively low concentration of LA, which produced effective analgesia with a wide margin of safety.

This study has certain limitations. First, SIFIB and CPB would require different positioning, so patients may not be blinded to the nerve block performed. The best way<sup>26</sup> to guarantee blinding would have been to perform the sham block and therapeutic block in each patient. We decided not to perform a sham block because we considered patients in category 3b of Serious Harm and Morbidity. 28 Second, spinal anesthesia may have the potential to last longer than six hours. However, we used a relatively low concentration of spinal anesthetic, and the median duration of analgesia would not have been expected to last more than 4.5 h.<sup>29</sup> In addition, Hu et al also chose dynamic pain score at 6

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h after THA as the primary outcome. <sup>13</sup> Thus, the time point of 6 h after surgery was deemed reasonable for the primary outcome. Third, we could not evaluate the real-time effects of CPB or SIFIB by checking the dermatomes for the blocklevel range because we administered the blocks after spinal anesthesia. However, all the block procedures were administered by experienced anesthesiologists who had performed CPB and SIFIB in more than 25 patients before the study. Therefore, we believe that all blocks have been implemented successfully. Finally, this was a single-centre study, and the findings warrant further investigation before application in general practice.

#### **Conclusion**

Ultrasound-guided CPB led to better postoperative pain relief, lower opioid consumption, and slightly affected muscle strength compared to SIFIB. Thus, CPB may be a promising approach for analgesia in patients undergoing THA.

## **Data Sharing Statement**

We all agree to share all the data for this article. The data that support the findings of this study are available from the corresponding author upon reasonable request.

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### **Disclosure**

The authors report no conflicts of interest in this work.

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