Upper Extremity Function after Transradial Percutaneous Coronary Intervention
Short-Term Interim Results of the ARCUS Study


1Department of Cardiology, Amphia Hospital, Breda, the Netherlands
2Department of Plastic, Reconstructive, and Hand Surgery, Albert Schweitzer Hospital, Dordrecht, the Netherlands
3Department of Cardiology, Albert Schweitzer Hospital, Dordrecht, the Netherlands
4Department of Cardiology, Erasmus Medical Centre, Rotterdam, the Netherlands
5Department of Cardiology, VU University Medical Centre, Amsterdam, the Netherlands
6Department of Plastic, Reconstructive, and Hand Surgery, Onze Lieve Vrouwe Gasthuis, Amsterdam, the Netherlands
7Department of Cardiology, Onze Lieve Vrouwe Gasthuis, Amsterdam, the Netherlands
8Department of Plastic, Reconstructive, and Hand Surgery, VU University Medical Centre, Amsterdam, the Netherlands
9Hand and Pols centre, Dordrecht, the Netherlands

#Authors contributed equally

*Corresponding author: Ijsselmuiden, A.J.J., MD, PhD, Department of Cardiology, Amphia Hospital, Mailbox 901584800 RK Breda, The Netherlands, Tel: 0031-0765954090; Fax: 0031-076-5953340; E-mail id: sijsselmuiden@amphia.nl

Abstract
Aims: To examine the short-term effects of transradial percutaneous coronary interventions on the upper extremities function.

Method and results: This is an interim analysis of the Effects of trAnsRadial perCUtaneous coronary intervention on upper extremity function (ARCUS) study. Out of 191 patients evaluated at 2 weeks after they had undergone a transradial percutaneous coronary intervention, 120 (62.8 %) had manifestations of upper extremity dysfunction on the side of the intervention. The main abnormalities were a decrease in sensibility, a ≥ 15 % increase in the Disabilities of the Hand and Shoulder questionnaire score and a ≥ 2 cm increase in hand and forearm volumes. Radial artery occlusions occurred in 12 patients in the upper extremity dysfunction versus 1 patient in the no upper extremity dysfunction group (p = 0.03). Patients with upper extremity dysfunction were significantly more likely to have a family history of heart disease (50 % vs. 26.8 %; p = 0.002). In addition, there was a trend (p = 0.07) toward a greater proportion of previous smokers in the group with upper extremity dysfunction than in the other group.

Conclusions: Upper extremity dysfunction after transradial percutaneous coronary intervention is a medical concern. However, our original score may be overly sensitive and overestimate the rate of upper extremity dysfunction. Further analyses are needed, as well as perhaps a modification of the primary endpoint.

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Keywords: Coronary angioplasty; Percutaneous coronary intervention; Transradial artery access; Upper extremity function; Radial artery occlusion

Abbreviations
BCTQ: Boston Carpal Tunnel Questionnaire; DASH: Disabilities of the Arm, Shoulder and Hand; PCI: Percutaneous Coronary Intervention; SD: Standard deviation; UED: Upper Extremity Dysfunction; VAS: Visual Analogue Scale

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Introduction

The transradial access is being increasingly adopted as the preferred approach for angioplasties, as it is associated with significantly lower rates of procedural complications, shorter hospitalisations, greater patient satisfaction and lower costs than the transfemoral access, which, on the other hand, is technically less challenging and associated with a steeper learning curve\(^\text{[1-5]}\). However, the precise effects of transradial procedures on the upper extremities function are unknown. The few published studies lack clear definitions of outcomes and systematic protocols and report inconsistent relationships between access-site complications and upper extremity function\(^\text{[6-9]}\).

Several important factors combined contribute to the normal function of the upper extremities, including intact anatomy, blood supply and lymphatic drainage, muscle strength, range of motion, coordination and sensibility. The absence of pain is a prerequisite for a normal upper extremity function, since pain has negative repercussions on these other factors. Upper extremity function has been defined as “The physiological capacity in which the patient can use an anatomically intact upper limb in everyday activities”\(^\text{[9]}\). Procedural complications may be the source of Upper Extremity Dysfunction (UED).

We undertook this prospective, observational study, to examine the effects of transradial Percutaneous Coronary Interventions (PCI) on the function of the upper extremities, and to optimize the transradial access-technique.

Methods

The aims and design of the “Effects of tranSRadial perCUtaneous (ARCUS) coronary intervention on upper extremity function” study have been published previously\(^\text{[4]}\). A target enrolment of 500 patients has been planned.

Diagnostic procedures

Doppler ultrasound examinations (SD2 Doppler, Huntleigh) were performed to examine both radial arteries before and after the PCI. Pain, disability of the upper extremities and the presence of carpal tunnel syndrome were ascertained, using self-administered questionnaires, including, respectively, the Visual Analogue Scale (VAS) for pain, the Disabilities of the Hand, Arm and Shoulder (DASH) questionnaire, and the Boston Carpal Tunnel Questionnaire (BCTQ). Swelling of the hands and forearms was measured by volumetry, using an ordinary tape-measure, and sensibility of the fingertips was measured with the Weinstein Enhanced Sensory Test (WEST filaments, Connecticut Bio instruments). We measured the key and palmar grip to ascertain the strength of the hands and thumbs. Additionally, we measured the isometric strength of flexion and extension of the hands and wrists. All measurements were performed on both upper extremities, according to the clinical assessment recommendations of the American Society of Hand Therapists\(^\text{[10]}\). All measurements were made pre procedural, at 24 hours, at 2 weeks, and at 1 and 6 months after the index procedure.

Percutaneous coronary intervention

The operators were strongly advised to adhere to the international professional practice guidelines for the performance of PCI\(^\text{[10]}\). However, procedural preferences were left to the discretion of each operator. After local anaesthesia of the wrist with a subcutaneous injection of 2% lidocaine, the radial artery was punctured with an introducer needle (Terumo Medical Corporation, Tokyo, Japan) using a modified Seldinger technique\(^\text{[12,13]}\). After introduction of the wire, the introducer needle was removed and a 6F introducer sheath (Terumo) was inserted\(^\text{[9]}\). Heparin, 100IU/KG and a mixture of verapamil, 5 mg, nitroglycerin, 200 μg, and 10 ml of saline solution were administered. A 6F hydrophilic guiding catheter (PendraCare, Leek, and the Netherlands) was inserted through the sheath and balloon angioplasty or stent implantation was performed in ≥ 1 coronary arteries. After completion of the procedure, the access-site was compressed using a transradial Band compression device (Terumo) injected with 13 cc air. After two hours the compression device was gradually deflated in 24 hours, according to the manufacturer’s protocol. The patient was provided with a sling and in case of swelling or haematoma of the arm with a minimal elastic compression bandage.

Study endpoints

The primary study endpoint is upper extremity function at two weeks compared to the pre procedural baseline measurements. Because of the absence of a precise, published definition and measurements of upper extremity function, hand experts tried to capture upper extremity function with the help of several examinations, performed according to the American Society of Hand Therapists\(^\text{[10]}\). The primary endpoint they composed is a binary score, absence or presence, of Upper Extremity Dysfunction (UED). Presence of UED is defined as a) a ≥ 1-point increase in the score of the BCTQ, or b) presence of ≥ 2 of the following criteria: 1) an increase in one of the questionnaire scores, 2) absence of radial artery signal on Doppler ultrasound examination, 3) ≥ 2 cm increase in forearm or hand volume, 4) a ≥ 1 filament decrease in fingertips sensibility, and 5) a decrease in wrist, elbow, key or palmar grip strength (Table 1)\(^\text{[11]}\). Secondary endpoints that are discussed in this paper are the complications following transradial PCI and the referral to a hand surgeon or rehabilitation specialist.

Table 1: Composed primary endpoint; positive binary score* of upper extremity dysfunction.

<table>
<thead>
<tr>
<th>Endpoint Description</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>≥ 1 point increase in either the symptom-severity score or the functional-status score of the BCTQ</td>
<td></td>
</tr>
<tr>
<td>Increase in the VAS score regarding the upper extremity of ≥ 2 points compared to baseline.</td>
<td></td>
</tr>
<tr>
<td>Absent signal of the radial artery during Doppler ultrasound examination.</td>
<td></td>
</tr>
<tr>
<td>≥ 60N decrease in palmar grip strength compared to baseline.</td>
<td></td>
</tr>
<tr>
<td>≥ 12N decrease in key grip strength compared to baseline.</td>
<td></td>
</tr>
<tr>
<td>≥ 15 % decrease in isometric strength compared to baseline.</td>
<td></td>
</tr>
<tr>
<td>≥ 1 filament increase in sensibility of the hand according to the WEST, compared to baseline.</td>
<td></td>
</tr>
<tr>
<td>≥ 1 cm increase at volumetry of the hand, using the figure-of-eight method, compared to baseline.</td>
<td></td>
</tr>
<tr>
<td>≥ 1 cm increase at volumetry of the forearm, measured circumferentially 8 cm distal of the medial epicondyle, compared to baseline.</td>
<td></td>
</tr>
</tbody>
</table>

\(^*\) Included were any endpoints that met the definition of Upper Extremity Dysfunction (UED).
If there are clinical complaints present at follow-up, patients will be referred to a hand rehabilitation specialist. Subsequently, diagnosis and diagnostic procedures performed at the hand centre will be registered. Also, the administered therapy and upper extremity related absence of work in days will be registered. These results will be published and analysed in the final paper.

Statistical Analyses

The details of the statistical analyses have been published previously. The results are presented as means ± SD and counts and percentages. Two groups of patients are distinguished: patients with or without upper extremity dysfunction according to protocol definitions. Between-group differences of dichotomous variables was analysed using Fisher’s exact test. Categorical variables with more than two categories are analysed with the Pearson chi-square test. For normally distributed variables the independent sample t-test was used, and the Mann-Whitney U test for continuous variables that are not normally distributed. Additionally, differences in the two upper extremity scores (intervention side – non-intervention side) collected in each patient will be analysed using Mc Nemar’s test. All tests were two-sided, with a statistical significance level set at 5%.

Interim results

Baseline characteristics: An interim analysis with 200 of the envisioned 500 patients was pre specified in the study protocol. Of these 200 patients 191 patients completed the pre procedural baseline and 2 week follow-up examinations of the upper extremity function in the context of the ARCUS study. The other 9 patients were either lost to follow-up or missed the 2 week follow-up visit. The baseline characteristics of the entire sample and of the groups with and without UED are shown in Table 2.

Table 2: Baseline characteristics of the entire study sample and of the groups with and without upper extremity dysfunction.

<table>
<thead>
<tr>
<th></th>
<th>All patients n = 191</th>
<th>Upper extremity dysfunction Present n = 120</th>
<th>Absent n = 71</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Men</strong></td>
<td>155 (81.2)</td>
<td>95 (79.2)</td>
<td>60 (84.5)</td>
</tr>
<tr>
<td><strong>Age, y</strong></td>
<td>64.8 ± 10.2</td>
<td>64.1 ± 10.6</td>
<td>66.1 ± 9.6</td>
</tr>
<tr>
<td><strong>Body mass index</strong></td>
<td>28.0 ± 4.5</td>
<td>28.1 ± 4.7</td>
<td>27.8 ± 4.3</td>
</tr>
<tr>
<td><strong>Height</strong></td>
<td>176 ± 9</td>
<td>176 ± 9</td>
<td>177 ± 9</td>
</tr>
<tr>
<td><strong>Smoking</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Current</td>
<td>32 (16.7)</td>
<td>21 (17.5)</td>
<td>11 (15.5)</td>
</tr>
<tr>
<td>Previous</td>
<td>94 (49.2)</td>
<td>53 (44.2)</td>
<td>41 (57.7)</td>
</tr>
<tr>
<td>Never</td>
<td>63 (32.9)</td>
<td>45 (37.5)</td>
<td>18 (25.4)</td>
</tr>
<tr>
<td><strong>Hypertension</strong></td>
<td>102 (53.4)</td>
<td>61 (50.8)</td>
<td>41 (57.7)</td>
</tr>
<tr>
<td><strong>Dyslipidaemia</strong></td>
<td>63 (32.9)</td>
<td>36 (30.0)</td>
<td>27 (38.0)</td>
</tr>
<tr>
<td><strong>Diabetes</strong></td>
<td>35 (18.3)</td>
<td>19 (15.3)</td>
<td>16 (23.9)</td>
</tr>
</tbody>
</table>

Values are means ± SD or numbers (%) of observations; *p = 0.002; all other between-groups differences are statistically non-significant

Group with upper extremity dysfunction

According to our binary score 120 of the 191 patients (62.8%) presented with dysfunction of the intervention hand at 2 weeks after the procedure. Figure 1 illustrates the various functions that were impaired. The main components of UED were a decrease in sensibility, observed in 41.7% of patients, a ≥ 15% increase in the DASH questionnaire score in 38.3% of patients, and an increase in hand volume in 38.3% of patients.

Complications

At two weeks, the radial artery was occluded in 12 patients (10%) in the UED group versus 1 patient (1.4%) in the group without UED (p = 0.003; table 3). By Doppler ultrasound
Modified upper extremity dysfunction of the non-intervention hand. Furthermore, some patients presenting with radial artery occlusions, the main components of UED were an increase in all the questionnaire scores, a decrease in sensibility and a decrease in strength of extension of the wrist.

### Clinical relevance

After two weeks of follow-up, 8 patients with UED (6.6%) were referred for rehabilitation. The main components of UED in these patients were an increase in pain with ≥ 2 points on the Visual Analogue Scale, an increase in DASH questionnaire and BCTQ scores, and a decrease in sensibility. The main complaint, which predominantly determined the patient referrals, was pain, often accompanied by a haematoma. The patients in the UED group referred for rehabilitation had an average of 5.7 ± 2.3 points, therefore, nearly 6 positive criteria.

Patients with UED were significantly more likely to present with a family history of heart disease (50% vs. 26.8%; p = 0.002). In addition, there was a trend (p = 0.07) toward a greater proportion of previous smokers in the UED group. In the group without dysfunction of the intervention hand, an asymptomatic music teacher was the only patient referred for rehabilitation and close surveillance after prolonged bleeding and development of a major haematoma at the access-site.

### Modified upper extremity dysfunction

Our primary study endpoint emphasises the questionnaire scores, the BCTQ score in particular. However, these questionnaire scores are not side-specific and overestimate the dysfunction of the non-intervention hand. Furthermore, some cut-off points may be overly stringent. Therefore, we adjusted other criteria. We used absolute numbers initially to compare the two-between groups differences are statistically non-significant.

### Table 3: Complications of transradial percutaneous coronary intervention.

<table>
<thead>
<tr>
<th>Upper extremity dysfunction</th>
<th>Present n = 120</th>
<th>Absent n = 71</th>
</tr>
</thead>
<tbody>
<tr>
<td>Minor bleeding at access site</td>
<td>0</td>
<td>1 (1.4)</td>
</tr>
<tr>
<td>Radial artery occlusion</td>
<td>12 (10.0)</td>
<td>1 (1.4)*</td>
</tr>
<tr>
<td>Access-site haematoma</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&gt; 5 cm</td>
<td>9 (7.5)</td>
<td>5 (7.0)</td>
</tr>
<tr>
<td>&lt; 5 cm</td>
<td>14 (11.6)</td>
<td>4 (5.6)</td>
</tr>
<tr>
<td>Swelling</td>
<td>7 (5.8)</td>
<td>1 (1.4)</td>
</tr>
<tr>
<td>Referral to hand rehabilita-</td>
<td>8 (6.6)</td>
<td>1 (1.4)</td>
</tr>
</tbody>
</table>

Values are means ± SD or numbers (%) of observations; *p = 0.003; all other between-groups differences are statistically non-significant.

### Positive UED score (dysfunction present): ≥ 2 criteria as specified below at the two-week follow-up:

- ≥ 2 points increase in visual analogue pain scale
- ≥ 2 points increase in Doppler signal
- ≥ 2 cm increase in volume of the hand
- ≤ 15% decrease in key pinch strength
- ≤ 15% decrease in palmar grip strength
- ≥ 2 filaments increase in sensitivity
- ≥ 2 cm increase in hand volume marginally significant, 8 cm distal of the medial epicondyle
- ≥ 2 filaments increase in sensitivity of the hand by the Weinstein Enhanced Sensory Test
- Decrease in key pinch strength compared to baseline.
- Decrease in palmar grip strength compared to baseline.
- Decrease in isotonic strength (flexion & extension) of the wrist and elbow.

### Discussion

This pre-specified interim analysis was conducted because this field of research is virtually a clean slate. There has been such an extended and all-round study grasping all facets forming upper extremity function after transradial PCI. The highly sensitive binary score, composed by hand specialists, is a first attempt to measure the overall function of the upper extremity after transradial PCI. We think that it may have been
too sensitive and overestimated the actual and clinically relevant upper extremity dysfunction. We tentatively hypothesise that we chose unforgiving cut-off values, which lead to high false positive UED scores in both hands. Therefore the primary endpoint should be modified, to optimize the specificity.

Other potential causes of a high UED score, either in the intervention arm or the non-intervention arm, might have been a pre-existent disease of the hands, such as arthrosis, carpal tunnel syndrome or tendinitis. In patients presenting with upper extremity diagnoses, the pinch strength of the asymptomatic side may decrease from compensating which results in overuse and fatigue. This must be taken into account when interpreting pinch strength of the asymptomatic extremity[14]. Furthermore, by increasing the peripheral arterial osmolality and causing vasodilatation[13], the contrast material may cause bilateral dysfunction. It may also injure the microvasculature and vascular endothelium by inducing apoptosis, and impair the physiological integrity resulting in UED, by altering vascular haemostasis, vessel growth, angiogenesis and permeability and tone of the blood vessel wall[16,17]. These disturbances may be observed with beta-adrenergic blockers and with statins, in the form of myalgia, myositis, myopathy, cramps and muscle weakness[18-20]. Diabetes may condition the prognosis of acute coronary syndromes and when regulated tight during the procedure it up-regulates the endothelial progenitor cell level and differentiation in patients going through an acute ST-elevation myocardial infarction[21,22]. We did not find this in our study.

Patients suffering from heart disease are hypoactive in the post procedural period, which might influence the questionnaire scores, and overestimate the degree of dysfunction. Furthermore, all questionnaire scores are not side-specific and could therefore overestimate the dysfunction on the intervention hand and non-intervention hand. We believe that in 39 patients (20%), the dysfunction in the non-intervention hand could be explained by an unspecific DASH score. Another 15% could be clarified by using absolute values of strength instead of percentages. Our presumption is that in another 10 - 15% it was induced by drugs.

The actual clinical relevance of our UED-score remains to be determined. We believe that the actual UED rate most likely will range between the rate of referrals for rehabilitation (6.6%) and the modified UED rate (16.9%). At the moment, the referral rate reflects the number of patients who had complaints and were urging for a referral to the hand rehabilitation specialist. However, during follow-up, several patients were candidates for referrals, though they were not all willing to see an additional physician and undergo further ambulatory clinic visits. They preferred to endure their disability, which caused an underestimation of the indications for referrals.

Keeping the complications and possible impairments in mind, custom treatments should be planned, according to professional and daily activities. Dialogs with the patients, for instance musicians, surgeons or interventional cardiologists are important, since the digital sensibility is one of the most often impaired functions, which could lower their professional performance.

In response to the results of this interim analysis, we already initiated a femoral control group. With this group, drug-induced complications could be demonstrated or excluded. Furthermore, the effects of pre-existent hand disease could be detected. Thus far, the non-intervention arm has been used as control. However, it is not accurate enough with respect to the questionnaires, since one needs both arms to open a bottle or to get dressed. Moreover, it would be worthwhile to focus on the prevention of UED by using slender and sheath less techniques, 4 or 5F catheters and miniaturization of the PCI equipment, and more efficient use of the closure devices, for example using oximetry-guided paten haemostasis[23,24]. Furthermore, the OPERA trial, which focuses on prevention and treatment, is being launched. Imaging of the radial lumen, using optical coherence tomography will provide more insight into the injury caused to the vessel by transradial PCI and into the pathogenesis of radial artery occlusion, thus in greater expertise of prevention and treatment of complications. Additionally, a study regarding treatment of complications, cost effectiveness and early referral is on-going.

Limitations of our study

The results of this interim analysis should be interpreted cautiously. At the completion of our enrolment, multiple variable analyses will be performed, presumably yielding more reliable results. Furthermore, this study was not randomised; the measurements were made in both upper extremities and the non-intervention extremity was used as the control arm. To eliminate the confusion caused by the high rate of dysfunction observed in the non-intervention hand, a femoral control group has been introduced. This should isolate the true effect of transradial PCI on the upper extremity function by eliminating the possible effects of drug-induced myopathies. It should also be stressed that these are short-term, interim results, while the long-term results are awaited to reveal the true dimensions and scope of these findings.

Conclusion

UED after transradial PCI is a true medical concern. However, our original score may be overly sensitive and overestimates the rate of UED. Further analyses are needed, as well as perhaps a modification of the primary endpoint. We expect the true clinical relevance to be between 6.6% - 16.9%.

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