

MISSION

CLINICAL REGISTRY

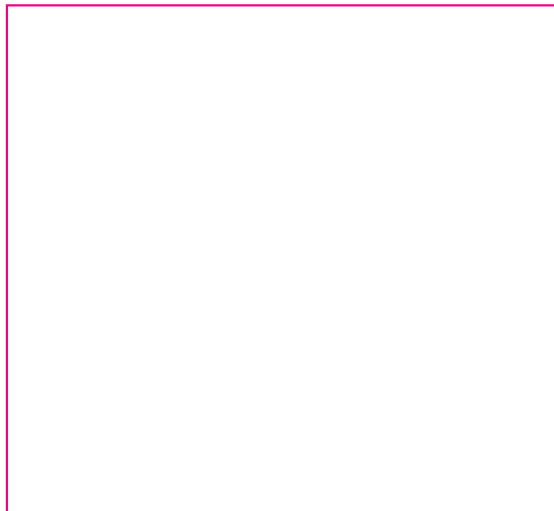
Common symptoms caused by carpal tunnel syndrome include:

- Numbness, tingling, burning, or pain in the thumb and index, middle, and ring fingers
- Occasional shock-like sensations that radiate to the thumb and index, middle, and ring fingers
- Pain or tingling that may travel up the forearm toward the shoulder
- Weakness in the hand
- These symptoms may keep you up at night

Why participate in the MISSION clinical registry?

- Surgical treatment may help alleviate your CTS. Please discuss with your doctor
- Carpal tunnel release procedures have been performed for almost 100 years
- Procedural approaches and techniques have improved over time
- The goal of any carpal tunnel surgery is to relieve pressure on the median nerve by dividing a ligament in your hand
- CTR-US is performed through a small (~ 5 mm) wrist incision
- The purpose of this clinical registry is to evaluate the long-term effectiveness of a minimally invasive technique: CTR-US using the FDA-cleared device, UltraGuideCTR™
- Data collected from the clinical registry will be used to educate healthcare providers and their patients about CTR-US as an option for carpal tunnel release

For more information,
please ask your doctor, visit
CTRstudy.com, or call 888-285-0404.



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Clinical Registry sponsored by Sonex Health, Inc.
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**Do you
experience pain
and numbness
in your hand?**

**Does this pain or
numbness wake you
up at night?**

**Do you find it difficult
to grip objects with
your hand?**

If so, you may be one of the millions of people suffering from a condition called **carpal tunnel syndrome**, which causes pain and tingling in your hand due to a compressed nerve.

NOW ENROLLING IN A CLINICAL REGISTRY

We are now enrolling in a clinical registry evaluating the effectiveness of a minimally invasive technique: carpal tunnel release with real-time ultrasound guidance (CTR-US) using the FDA-cleared device, UltraGuideCTR™.

For more information, please visit CTRstudy.com or call 888-285-0404.



CTR with Ultrasound Guidance

Who can join the clinical registry?

- You are ≥ 18 years of age
- You have a clinical diagnosis of carpal tunnel syndrome in one or both hands
- Your symptoms have not improved with other treatment options including activity modification, bracing, splinting or injections
- Your physician considers you a surgical candidate
- You agree to complete periodic follow-up questionnaires over a 2-year period
- You have a valid smartphone number and/or email address to receive and answer follow-up questionnaires

What is a clinical registry?

A clinical registry is a database that collects information about your health and the care you receive as a patient. The data in the clinical registry comes from information you provide about your care before and after your procedure and is added to information on other patients who are similar to you. It is then used to help improve the quality of your care as well as the care of other patients, now and in the future. In this clinical registry, the research is evaluating the long-term effectiveness of CTR-US.

Who is conducting this clinical registry?

- This clinical registry is led by a physician (principal investigator) and includes physicians from a number of clinical sites across the United States
- This clinical trial is sponsored by Sonex Health, Inc. under an IRB-approved study number 1360397 and ClinicalTrials.gov Identifier NCT06071468

How does this clinical registry work?

- Your physician will evaluate and discuss with you whether surgery is indicated to help the problem
- If surgery is indicated, your physician will discuss whether you are eligible for this registry
- A series of tests and questionnaires will be given to you to verify if you meet the criteria to be a part of the registry. This is called screening and will be done before you qualify to receive treatment as a part of the registry
- If you qualify to be in the registry, you will be scheduled for CTR-US

What type of follow-up is required?

If you decide to join and qualify for this clinical registry, you will be expected to complete online follow-up questionnaires through a secure website at 1-7 days, 2 weeks, and 1, 3, 6, 12, and 24 months after the procedure.

Will I be paid to join the clinical registry?

If you qualify for the clinical registry, you will be paid up to \$600 to share your experiences at 7 follow-up time points after the procedure.

How is CTR-US different from other carpal tunnel release techniques?

Carpal tunnel release surgical procedures have been performed for almost 100 years. Over time, procedures and technology have improved to help reduce the invasiveness of these traditional surgical procedures. Today, several techniques are utilized, including traditional open, mini-open, endoscopic, and ultrasound-guided. Incision sizes and locations may vary, but the goal of each technique is to relieve pressure on the nerve by dividing the transverse carpal ligament.

- CTR-US uses a minimally invasive technique in which a small incision is made in the wrist and the physician sees the carpal tunnel anatomy through ultrasound
- CTR-US is typically performed under local anesthesia

Am I required to be in the clinical registry?

No. Your participation is completely voluntary. If you enroll in the clinical registry, you can still leave the registry at any time without having to provide a reason.

How long will I be in the clinical registry?

You will be expected to complete online follow-up questionnaires through a secure website at 1-7 days, 2 weeks, and 1, 3, 6, 12 and 24 months after the procedure.