

TINEARITY™ G1 PATIENT GUIDANCE

Tinearity™ G1 patient guidance

There exist two ways by which a patient may ascertain the appropriateness of using the Tinearity G1 for the treatment of their tinnitus condition:

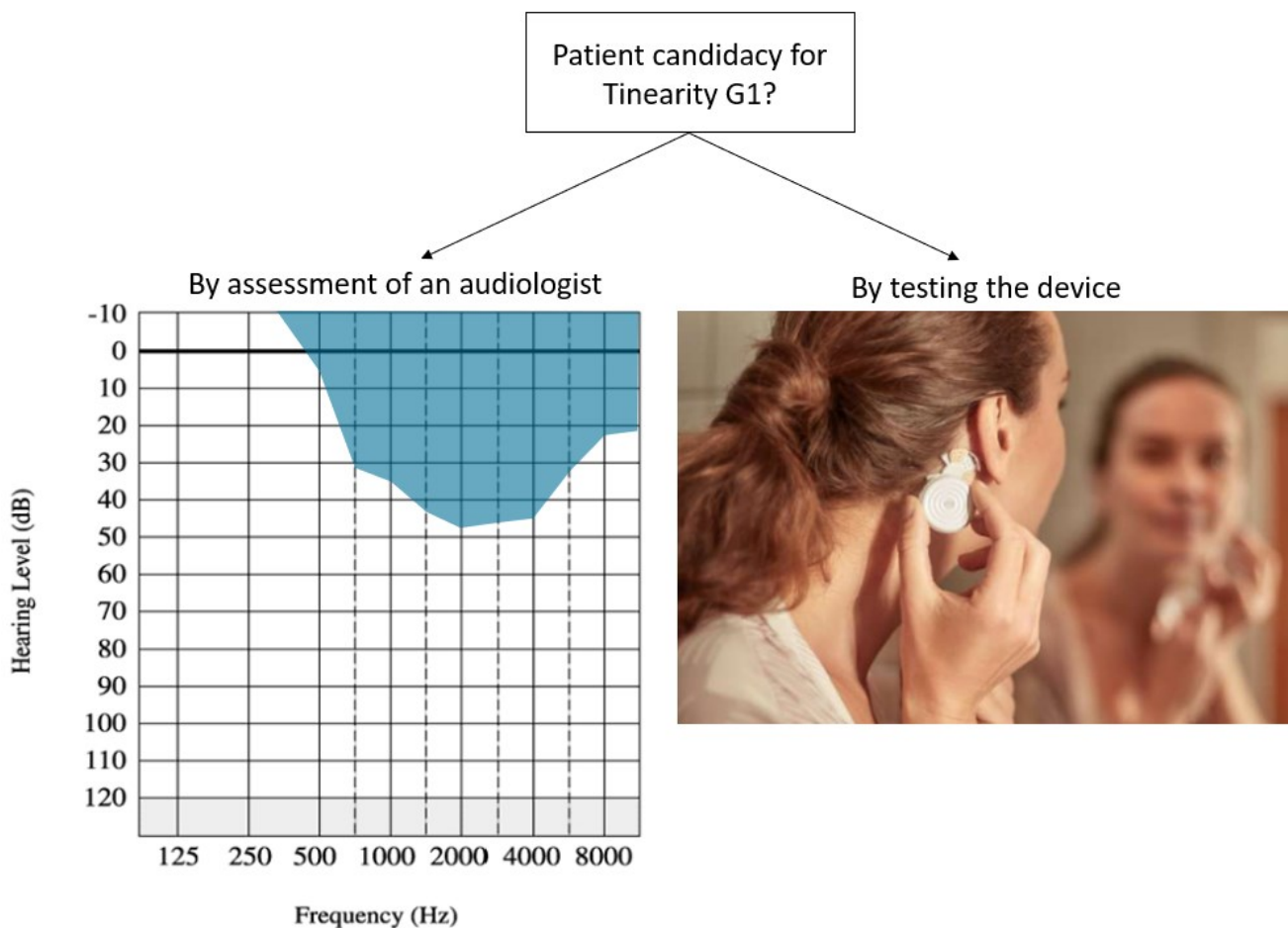
1) Through audiology assessment

Engage the services of a qualified audiologist who will conduct an assessment analysis of the frequency and amplitude of your tinnitus auditory sensation and to assess if you have normal hearing. Then evaluate if the frequency and amplitude falls within the indication area of the Tinearity G1 device illustrated in the blue shaded area in the figure below. If it does and if you have normal hearing then the Tinearity G1 device is suitable for Tinnitus relief treatment.

2) By direct device testing

Simply connect the Tinearity G1 device, according to the user manual behind your ear on the same side as where the tinnitus is. If bi-lateral, start at the same side as where the tinnitus is worst. Gradually amplify the device's output volume until the audible perception of tinnitus becomes imperceptible. Apply the second sound generator behind the other ear and adjust the volume to perceive a homogenic sound. If you can mask the tinnitus sound before you reach maximum volume of the device then the Tinearity G1 device is suitable for Tinnitus relief treatment.

Figure to the left shows the indication area (blue shaded) of Tinearity G1 in an audiogram.



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