Project Design Specifications
Tactile Auditory Sensory Substitution
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Function:

High frequency hearing loss is the most common form of hearing loss experienced. It is caused by damaged nerve ends on the hairs in the cochlea. People with high frequency hearing loss cannot hear high frequency sounds such as ‘s’, ‘f’, ‘sh’, ‘ch’, ‘t’, and ‘p’ sounds. Since these sounds are some of the most used in the English language, high frequency hearing loss is a major problem that inhibits communication immensely.

Individuals with high frequency hearing loss tend to rely on hearing aids, which amplify sounds. However, the hearing loss cannot be fixed with amplification of these high frequency consonants because the person simply cannot hear in that frequency range, the volume makes no difference. Instead of amplification, these missing consonants can be communicated by sensory substitution. In other words, a sense other than hearing can be used to send the auditory information to the brain. The goal is to design and develop an auditory substitution device that through the use of vibro-tactile stimulation can substitute for regional frequency hearing loss. Individual vibrators will represent certain frequency ranges, and when the user feels the specific vibration attributed to a certain frequency range, the user will be able to identify the sound.

After using the device for an extended period of time, the user will no longer have to concentrate on the vibrations but will be able to “hear” the high frequency sounds. Due to the plasticity of the brain, the brain will interpret the vibrations as sounds and fill in the gaps for the user, and communication will be restored.

Client Requirements:
- The device will substitute for high frequency hearing loss to the extent of helping the user in everyday communication.
- The device will use vibro-tactile stimulation.
- The device should be self contained, portable, and discrete.
- The device should not cause discomfort to the user.
- A testable prototype will be completed for analysis of frequency range and wearability.

Design Requirements:

1. Physical and operational characteristics
   a. Performance requirements
      - It will increase the user’s quality of communication by allowing the user to recognize high frequency consonants and incorporate them into word recognition through vibro-tactile stimulation.
• This device should use the programmable functions of computer software to recognize and separate certain high frequency sounds and communicate them to the vibro-tactile stimulator.

b. Safety
• A current of more than 5 mA should not pass through the device and into the user.
• The device should not heat to over 43° C (110° F) while in use.

c. Accuracy and Reliability
• The device should be able to pick up sound that is detected at normal speaking level of 60 dB, and transmit vibrations according to the frequency of the particular sound.
• The device should be able to process and substitute for the consonants T, F, S, Th, Sh, and P when coming from a variety of different vocal tones. It should be accurate enough to improve scores on standard speech recognition evaluation tests, Word Intelligibility by Picture Identification (WIPI) tests, sentence level Bamford-Kewal-Bench tests (BKB), and PLOTT tests for vowel and consonant discrimination.
• Human Hearing Frequency Range: 20 – 20,000 Hz
• Speech Frequency Range: 125 – 8,000 Hz
• High Frequency Hearing Loss: above 1,000 Hz

<table>
<thead>
<tr>
<th>Sound</th>
<th>Frequency (Hertz)</th>
</tr>
</thead>
<tbody>
<tr>
<td>T</td>
<td>3500</td>
</tr>
<tr>
<td>F</td>
<td>4000</td>
</tr>
<tr>
<td>S</td>
<td>4000</td>
</tr>
<tr>
<td>Th</td>
<td>4000</td>
</tr>
<tr>
<td>Sh</td>
<td>2000</td>
</tr>
<tr>
<td>P</td>
<td>1500</td>
</tr>
</tbody>
</table>

d. Life in Service
• The transducers should last at least 3 years.
• The sound processing unit, along with its microphone, should last 10 years.
• Adhesive should last at least through an entire day.
• On a single battery charge the device should last approximately 5 days, similar to that of a common hearing aid so they can be charged at the same time.
• Common hearing aid batteries have an output voltage of 1.4 V and have power ratings between 140 and 640 mAh. With daily use of the device being about 14 hours the device should draw from 2 - 10 mA of current from the battery.

e. Operating Environment
• The device will be located near the ear.
• Elements such as wind, rain, sun and sweat should not cause the device to vibrate for non-spoken noises and should not cause the device to output dangerous levels of current or distort outgoing signals.
• Eventually test the prototype on human subjects in a lab setting.

f. Ergonomics
• The device should not move during normal physical activity.
• The transducer unit should be hidden by the ear.
• The device should be comfortable to the user.
• The controls of the processing unit should be small enough to keep them discrete yet still be able to be manipulated.
• The vibro-tactile device will be attached with adhesive that will not cause irritation.
• The location of the transducer should be easily repeatable.

g. Size
• The transducer unit should be no more than 5 cm in length, 1.75 cm wide, and 1.25 cm thick.
• The processing unit should be no more than 10 cm in length, 5 cm wide, and 2 cm thick.

h. Weight
• The weight of the processing unit should be no more than 8 oz.

i. Materials
• Adhesive which holds the transducer unit in place should not irritate skin, leave large amounts of residue, or be painful to remove.
• Soft, durable plastic such as vinyl should be used.

j. Aesthetics, Appearance, and Finish
• Unit should be flesh-colored and not overtly noticeable to others.
• Adhesive attachment used for transducer unit should not leave large amounts of residue and should not be painful to remove.

2. Production Characteristics

a. Quantity
• The device should be able to be produced in mass quantities.

b. Target Product Cost
• The device should cost between $300 and $500.
• This price is approximately 5-10% of the total cost of a hearing aid.

3. Miscellaneous

a. Standards and Specifications
• FDA approval of a class II device.
• Must follow regulation codes 21 CFR 812 and 21 CFR 50 for testing human subjects.

b. Customer
• The device should have adjustable frequency ranges depending on what the user needs most or the environment the device is operating in.

c. Patient-related concerns
• The device should not cause discomfort in any way.
• The device should not be overly noticeable.

d. Competition:
• Tickle Talker
• Tacticon 1600
• Tactaid 7
  o http://www.tactaid.com/tactaid71.html