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Functional near-infrared spectroscopy for neuroimaging in cochlear implant recipients

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#### **Functional near-infrared spectroscopy for neuroimaging in cochlear** 1

## **implant recipients[1](#page-1-0)**

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#### **Abstract** 29 28

<span id="page-1-0"></span>1Abbreviations: BOLD, blood-oxygen level dependent; CI, cochlear implant; CW, 2 3 continuous wave; EEG, electroencephalography; FD, frequency-domain; fMRI, functional magnetic resonance imaging; fNIRS, functional near-infrared spectroscopy; 4 5HbO, Oxygenated hemoglobin; HbR, Deoxygenated hemoglobin; MEG, 6magnetoencephalography; NIR: near-infrared; PET, positron emission tomography; TD, 7time-domain; SNR, signal-to-noise ratio; SRT, speech reception threshold

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31 Functional neuroimaging can provide insight into the neurobiological factors that 32 contribute to the variations in individual hearing outcomes following cochlear 33 implantation. To date, measuring neural activity within the auditory cortex of cochlear 34implant (CI) recipients has been challenging, primarily because the use of traditional 35 neuroimaging techniques is limited in people with CIs. Functional near-infrared 36 spectroscopy (fNIRS) is an emerging technology that offers benefits in this population 37because it is non-invasive, compatible with CI devices, and not subject to electrical 38 artifacts. However, there are important considerations to be made when using fNIRS to 39 maximize the signal to noise ratio and to best identify meaningful cortical responses. This 40 review considers these issues, the current data, and future directions for using fNIRS as a 41 clinical application in individuals with CIs.

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43Keywords: fNIRS, cochlear implant, hearing loss, neuroimaging, speech

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### **471.** Introduction

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49 Cochlear implants (CI) have restored hearing to over 90,000 individuals in the United 50States in the past 30 years (FDA, 2015). Significant advances in speech processor design, 51 signal processing and surgical techniques have resulted in progressively enhanced 52 performance (Rubinstein, 2004; Roland et al., 2006; Srinivasan et al., 2013). As a result, 53 cochlear implantation has become a highly successful prosthetic solution to replace the 54 function of a sensory organ. Intervention with deaf children has been particularly 55 successful: many children who would otherwise have been placed in schools for the deaf 56 and taught sign language are now learning alongside mainstream peers in a regular 57 classroom environment. The primary goal of cochlear implantation is now open-set 58 auditory-only speech understanding in everyday listening environments. However, while 59the majority of implant recipients achieve this goal, many still perform poorly (Lazard et 60al., 2012; Miyamoto et al., 1994).

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62 The factors that contribute to the wide variations in individual outcomes following 63 cochlear implantation are diverse and not completely understood (Lazard et al., 2014; 64Peterson et al., 2010). Numerous reports have identified age of implantation as a strong 65 predictor of better CI outcome (e.g., the younger, the better) (Kirk et al., 2002; 66Nikolopoulos et al., 1999; Robinshaw, 1995). Investigators have also demonstrated that 67 children who communicate orally achieve better speech perception skills than children 68who use visual sign communication (Osberger and Fisher, 2000; Geers et al., 2003). 69 Finally, family income predicted language outcomes in pediatric CI recipients (Holt and

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70Svirsky, 2008). In order to more fully understand how such neurobiological, cognitive, 71 and societal factors influence language outcomes post-implantation, it may be beneficial 72to examine the neural processing during the perception of auditory stimuli through a 73 cochlear implant. Together with behavioral measures, neurophysiological indicators have 74the potential to guide post-implant programming in support of deaf patients' speech and 75 language outcomes and, eventually, even predict results for an individual CI patient 76before implantation occurs.

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78 Functional near-infrared spectroscopy has already been shown to be a reliable 79 neuroimaging modality in both adult and pediatric populations (Fava et al., 2014; Giraud 80et al., 2001; Quaresima et al., 2012; Wilcox, et al, 2005). Generally, reviews of this 81literature have focused on the use of fNIRS in research on language development and 82language processing in healthy populations (Crosson et al., 2010; Elwell and C. E. 83 Cooper, 2011; Gervain et al., 2011; Lloyd-Fox et al., 2010; Quaresima et al., 2012, Fava 84et al. 2011; Fava et al., 2014; Wilcox, et al, 2005). More recently, an emerging body of 85 reviews addresses the imaging instrumentation and methodology, as well as approaches 86to statistical analysis of fNIRS data (Bandettini, 2009; Piper et al., 2014; Scholkmann et 87al., 2014; Tak and Ye, 2014). However, most relevant to CI research is the fact that fNIRS 88 is compatible with these devices. This review explores applications and limitations of 89fNIRS in the CI population, comparing it with traditional neuroimaging methods. We 90 summarize the existing literature on the use of fNIRS in adult and pediatric CI recipients, 91 and conclude by outlining possible directions for future research and clinical applications 92using this promising imaging technique in the CI population.

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#### 942. Neuroimaging options in cochlear implant users

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96 Because auditory perception occurs within and beyond the auditory cortex, neuroimaging 97has the potential to provide an additional clinical measure for assessing whether the 98 electrical stimulation of the cochlea by the CI is reaching and stimulating auditory-99 specific cortical regions of the brain similar to normal-hearing subjects (Pasley et al., 2012; Steinschneider et al., 2014). Such information can supplement behavioral tests, 100 101 which are often limited in young CI users (Choi and Oghalai, 2005; Katzenstein et al., 2009; Lin et al., 2010; Oghalai et al., 2009; Santa Maria and Oghalai, 2014; Williamson 102 103et al., 2009; Ying et al., 2013). However, there are inherent limitations in the use of all of 104the currently available neuroimaging modalities in CI recipients, as outlined below and 105 summarized in Table 1.

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107 Functional neuroimaging attempts to identify the brain systems responsible for different 108behaviors by comparing brain activity during contrasting states (Aine, 1995; Crosson et 109al., 2010). The logic is that neurons in different areas of the brain associated with specific 110 cognitive processing tasks generate electrical signals when they are active. As a result of 111this activation, the metabolic needs of neurons change: increased oxygen demand results 112in increased cerebral blood flow and thus oxygen delivery to that area, with a consequent 113 decrease in deoxygenated hemoglobin (HbR) (Babiloni et al., 2009). Certain 114 neuroimaging modalities, such as EEG, measure this neural activation directly by 115 recording the average electric field potential at different regions of the scalp. In contrast,

116 metabolic neuroimaging methods, such as fMRI, PET, and fNIRS, are indirect, surrogate 117 measures of neuronal activity (Castañeda-Villa et al., 2012; Girouard, 2006; Levitin and 118Menon, 2005; Mc Laughlin et al., 2013).

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120Although functional neuroimaging technologies have the potential to provide insight into 121 the cortical changes that take place in patients with cochlear implants, obtaining 122 meaningful measurements of cortical responses in CI recipients has proven challenging. 123This is primarily because the traditional imaging methods have limitations when used in 124implanted patients, and so alternative neuroimaging strategies have been sought. In this 125 context, functional near-infrared spectroscopy (fNIRS) has been a welcome addition to a 126limited choice of neuroimaging modalities suitable for use in CI recipients. Here we 127 outline the primary techniques and assess their appropriateness for use in combination 128 with CIs. Because it is important understand the benefits and downsides to each 129technique when selecting an imaging modality, we briefly review several commonly-used 130techniques including fMRI, PET, EEG, and MEG, before moving on to an in depth 131explanation of fNIRS.

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#### 1332.1 Functional MRI

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135 Functional MRI provides high spatial resolution and is often the neuroimaging 136technology of choice in unimplanted subjects. However, conventional CIs are 137 incompatible with fMRI for several reasons. The primary reason is that CIs contain 138 internal magnets and ferromagnetic components, including a coil used to

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139transcutaneously relay data from the external processor to the surgically implanted 140 components (Doucet et al., 2006; Gilley et al., 2008; Majdani et al., 2008). Such 141 ferromagnetic implants exposed to electromagnetic fields or radiofrequency energy may 142heat, induce a current, or become dislocated (Azadarmaki et al., 2014; Portnoy and 143 Mattucci, 1991; Teissl et al., 1999). Thus, the most important concern in using fMRI to 144study a subject with a CI is patient safety. Furthermore, the magnet and coil interact with 145the electromagnetic fields found in MRI scanners, producing interference that can disturb 146 data transfer, and malfunction of the implant can occur due to demagnetization of the CI 147internal magnet via the imaging magnet (Majdani et al., 2008; Ponton et al., 2000). 148 Finally, CIs produce considerable artifacts on the MR image, obscuring cortical regions 149 proximal to the internal magnet (Majdani et al., 2009). Thus, these signal-void areas can 150 compromise accurate diagnosis of certain medical conditions when used for medical 151 imaging and make it nearly impossible to measure activity within the ipsilateral temporal 152lobe when used for functional imaging.

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154In response to these limitations, certain manufacturers have designed CIs with removable 155 internal magnets. Unfortunately, large artifacts often remain on the MRI even after the 156 internal magnet is removed (Risi et al., 2004). Other models of CI have MRI-conditional 157 internal magnets that do not need to be removed prior to scanning. Regardless of the 158 status of the internal magnet, the external processors for all CI devices are MRI unsafe 159(Azadarmaki et al., 2014) and the radiofrequency fields generated by the MRI interfere 160 with the transcutaneous radiofrequency link between the external and internal coils 161(Lazeyras et al., 2002; Seghier et al., 2005). Auditory stimulation by the implant during

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162 imaging is therefore generally precluded, though anatomical images can be acquired for 163 medical purposes (Baumgartner et al., 2001; Crane et al., 2010; Gubbels and 164McMenomey, 2006).

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166The limitations of using fMRI with the CI population extend beyond equipment 167 incompatibility issues. MRI is subject to movement artifacts (Quaresima et al., 2012), 168 requiring subjects to remain completely still and to avoid overt vocalizations while in the 169 scanner. In infants, this translates into the need for restraints and even sedation and/or 170anesthesia. Sedatives and anesthetics, of course, alter brain activity and therefore change 171 cortical responses to auditory stimuli (Marcar et al., 2006). Such circumstances 172 considerably restrict the use of fMRI in this age group.

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174It is also important to consider that fMRI is a noisy imaging modality, which introduces a 175 potential confounding effect as the background noise cannot be matched between deaf 176 and hearing participants (Dewey and Hartley, 2015). Moreover, the acoustic noise 177 associated with fMRI creates an intrusive testing environment for younger children and 178 disturbs the presentation of auditory stimuli relevant to CI users (Gervain et al., 2011). 179 Finally, the BOLD (Blood Oxygenation Level Dependent) signals obtained using fMRI 180 relate to changes in HbR only and do not directly convey information about HbO.

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1822.2 PET scan

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184Nuclear functional imaging techniques such as PET scans have more frequently been 185used in studies involving CI users. Previous investigators employed PET scans to

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186 examine various auditory cognitive processes in the CI population (Limb et al., 2010; 187 Naito et al., 2000; Wong et al., 1999), and several dedicated reports have even been 188 published for reviewing the use of PET scans in language processing research on CI 189 recipients (Aggarwal and Green, 2012; Giraud et al., 2001). Several factors account for 190the popularity of this neuroimaging modality for use with CIs among the scientific 191 community. First, PET is fully compatible with CIs. It also has good spatial resolution 192 and, as with MRI, it can image activity in deep, subcortical structures (Bandettini, 2009). 193Because PET is a relatively quiet imaging modality, it is suitable for studies involving 194 auditory stimuli. Finally, it is tolerant to subtle subject movements thanks to rapid image 195 acquisition times, a significant advantage over fMRI (Crosson et al., 2011).

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197The significant drawback of using this imaging modality is the exposure of the research 198 subjects to radiation and the necessary limitation in the number of scans that this implies. 199The radioactive tracers or carrier substances need to be injected into the blood stream, 200 which many subjects find aversive. For these reasons, PET is rarely used in research 201 studies involving children. Though understandable, this is unfortunate because children 202are a demographically important age group within the CI population. The use of PET to 203study neuroplasticity post-implantation is also ethically challenging, as measuring such 204 changes would require sequential longitudinal testing in the same subject (Giraud et al., 2052001). Limited temporal resolution, or the accuracy on a temporal scale with which a 206 neural event can be characterized (Crosson et al., 2010), is another shortcoming of PET. 207This is because PET's ability to resolve neural events is on the order of tens of seconds 208 compared to only a few seconds for fMRI (Bandettini, 2009). Such limited temporal

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209 resolution requires averaging over long blocks of events; higher sampling rates are 210 generally preferred in functional studies because they allow the use of event-related 211 paradigms, which offer greater flexibility and more precision in experimental inquiry 212(Aine, 1995).

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#### 2142.3 EEG and MEG

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216Unlike fMRI and PET, EEG and MEG directly measure the electrophysiological response 217 of neural activation. The resulting advantage of this technique is an unrivaled temporal 218 resolution in the sub-millisecond range (Babiloni et al., 2009), however at the expense of 219 spatial resolution (Posner and Levitin, 1997). Studies have shown that auditory evoked 220 potentials recorded in EEG provide a useful objective metric of performance in CI 221 patients (Castañeda-Villa et al., 2012; Mc Laughlin et al., 2013). It is therefore not 222 surprising that the EEG literature in CI users is abundant and, indeed, has greatly 223 contributed to the understanding of auditory processing in this population (Sandmann et 224al., 2010; Zhang et al., 2010). In addition, the combination of the high temporal 225 resolution and an excellent safety profile make EEG and MEG ideally suited for follow-226up studies requiring several successive assessments, such as those investigating cortical 227 plasticity following implantation (Doucet et al., 2006; Gilley et al., 2008). Finally, EEG is 228 tolerant to subtle movements and can even be used with fully awake infants.

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230On the other hand, as mentioned, EEG and MEG offer relatively poor spatial resolution 231 due to the inverse Poisson problem: the location of activity within a sphere is ambiguous

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232when measuring from the surface of that sphere (Posner and Levitin, 1997). While the 233 reconstruction of brain responses to specific cortical regions is possible (Ferree et al., 2342001; Song et al, 2015), the accuracy of this localization remains inferior to other 235 modalities such as fMRI or PET (Ponton et al., 2000). Data corruption by the electrical 236 components of the implant is another major limiting factor for the use of EEG in 237 combination with CIs. To minimize the electrical artifacts produced in EEG recordings, 238 only short auditory stimuli such as tone bursts or clicks can be employed in CI studies, 239 which significantly limits the flexibility of the experimental paradigm (Gilley et al., 2402008). Despite the various techniques that have been described to filter this artifact, the 241interpretation of auditory evoked potentials in EEG remains challenging (Mc Laughlin et 242al., 2013; Sandmann et al., 2009). Additionally, MEG measures very weak magnetic 243 fields that can only be recorded in magnetically shielded rooms equipped with detectors 244that are highly sensitive to minute changes in magnetic signals (Crosson et al., 2010). 245Similar to fMRI, MEG instrumentation interacts with the internal magnet of most CI 246 models, precluding any useful recording. To successfully monitor neural activity in CI 247users using MEG, certain conditions must be fulfilled. This unique experimental setup is 248 described by Pantev (Pantev et al., 2006), who reported the only MEG study involving CI 249users. The basis for the methodological success of this study is twofold. First, the two 250 participants enrolled were recipients of Clarion (Advanced Bionics, Valencia, CA) 251 magnet-less implants – now withdrawn from the market. Second, a unique radio 252 frequency shield was applied between the head of the patients and the MEG device, 253 preventing interference from radio frequency signals transmitted by the CI. Such setups, 254 however, are very rare and extremely costly.

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#### **3. fNIRS** 256

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258Before fNIRS was adapted for use in people with CIs, PET was reported to be the only 259technique suitable for measuring brain responses in the CI population for all of the 260 reasons outlined above (Giraud et al., 2001; Truy, 1999). Because the concepts, features, 261 and instrumentation of fNIRS have been described in substantial detail in previous 262 reports (Elwell and C. E. Cooper, 2011; Gervain et al., 2011; Lloyd-Fox et al., 2010; 263 Quaresima et al., 2012), we will only briefly address them in this review. Here we focus 264 primarily on the characteristics of fNIRS that are relevant to its use with the CI 265population.

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2673.1 General principles

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269fNIRS is an optical imaging technique: it uses near-infrared (NIR) light to detect changes 270in cerebral blood flow as a proxy for neural activation. When a beam of light is directed 271 onto tissue, three factors can interfere with its undisturbed propagation (i.e. transmission) 272through it: reflection/refraction, absorption and scattering (Niemz, 2002). The 273 contribution of reflection/refraction can essentially be ignored in opaque media such as 274the skull. The intensity of the transmitted light therefore depends on the amount of non-275absorbed and non-scattered photons(Welch, 2011; Gervain et al., 2011). Biological 276 tissues preferentially absorb light in the visible spectrum, while being relatively 277transparent to light in the NIR wavelengths (650-1000 nm) (Smith, 2011). As a result,

278NIR light can penetrate through superficial biological layers, enabling sampling of deeper 279 tissue structures. For neuroimaging, this means that fNIRS can effectively probe the 280 surface of an adult brain to a depth of up to 1.5 cm (Elwell and C. E. Cooper, 2011). 281

282fNIRS is capable of measuring changes in cerebral blood flow because hemoglobin is the 283 main pigmented molecule in human tissues that is present in clinically significant 284 quantities to exhibit oxygenation-dependent absorption of light in the NIR spectrum 285(Delpy and Cope, 1997). In tissues, hemoglobin exists in an oxidized (oxygenated 286hemoglobin, HbO) and reduced (HbR) form, each characterized by a unique absorption 287 spectrum. The aim of NIRS neuroimaging is to quantify the concentrations of these two 288hemoglobin chromophores in the tissues traversed by NIR light. This is possible using the  $289$ Beer-Lambert Law, an equation that describes the light absorbance  $(A)$  at a given 290wavelength  $(\lambda)$  in a medium (Crosson et al., 2010):

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$$
A=-\log\left(\frac{I}{I_o}\right)=c.\varepsilon_{\lambda}.
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292Shining light of an appropriate wavelength at a given intensity (incident light, I) on the 293head, and measuring the intensity of the light that leaves the tissues (transmitted light,  $I_0$ ) 294 allows for the calculation of the concentration of the medium, "c" (i.e. the concentration 295 of HbR, HbO and total hemoglobin). This concept assumes that the molar extinction 296 coefficient of the medium at that specific wavelenght  $(\epsilon_{\lambda})$  and the optical pathlength "*l*" 297 in the tissues (the path the light travels between the source and the detector) are known. 298

299The application of this physical principle forms the basis of fNIRS neuroimaging. Of 300 course, other factors need to be considered. Light scattering caused by skin, hair and

301 skull, also contributes to light attenuation in tissues, resulting in an unknown light loss 302that needs to be accounted for (Delpy and Cope, 1997). Furthermore, light does not travel 303through biological tissue in a straight line. The Beer-Lambert Law was therefore 304 modified to take into account the scatter and the non-linear trajectory of light in tissues, 305 referred to as the differential pathlength factor (Cope et al., 1988). These two factors 306 cannot be measured directly using continuous-wave NIRS systems (see below), therefore 307 only changes in HbO and HbR concentrations, as opposed to absolute values, can be 308obtained. A detailed description of the mathematical model underlying light absorption in 309 scattering media can be found elsewhere (Gervain et al., 2011; Hoshi, 2003; Sassaroli and 310 Fantini, 2004).

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312Practically speaking, fNIRS is performed on human subjects by placing a light source and 313a light detector adjacent to each other above the brain area to be measured. This source-314 detector pair is called a channel. A convex banana-shaped tissue region is sampled, 315 corresponding to the light path through the tissue between the source and detector. The 316 depth of penetration of the NIR light in brain tissue is approximately half of the source-317 detector distance. To reach a clinically relevant depth of cortical area, the source-detector 318 distance should be 2-3 cm in infants and 3-5 cm in adults (Quaresima et al., 2012). The 319 choice of the wavelength pair is also important, as it affects the quality of the fNIRS 320 signals. Ideally, one wavelength should be sensitive to HbO; the other to HbR. This is 321 possible because HbO and HbR demonstrate differential absorption in the NIR spectral 322range (except at the isosbestic point, where the extinction coefficients of these two 323 chromophores are equal). Generally, wavelengths below the isosbestic point are used to

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324 measure HbR responses (below 760–770 nm), whereas longer wavelengths are more 325 sensitive to HbO (up to 920 nm) (Boas et al., 2004). Theoretical models also revealed that 326the highest signal-to-noise ratios were obtained if one wavelength was below 720 nm, 327 and the other higher than 730 nm (Uludag et al., 2004). The 690 nm and 830 nm pair is 328 commonly reported in fNIRS literature, but a variety of other systems capitalizing on 329 different wavelength contrasts are commercially available (Lloyd-Fox et al., 2010).

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331 Three different fNIRS instrumentation techniques are currently available, and they vary 332in the type of illumination employed (Ferrari and Quaresima, 2012). The first modality, 333 continuous wave (CW) light, is the most commonly used and the least costly. It is based 334on constant tissue illumination and simply measures changes in light attenuation as it 335 passes through the head. This technique does not allow calculation of light scattering or 336 optical path length in tissues and, as a result, can only determine relative changes in HbO, 337HbR and total hemoglobin concentrations (Scholkmann et al., 2014). However, relative 338 values of hemodynamic parameters are usually sufficient in functional brain studies. The 339last two techniques, time-domain (TD) and frequency-domain (FD), are equivalent in that 340they both measure the time needed by light to travel through tissues (i.e. time of flight) to 341 determine optical path length (Wolf et al., 2007). They differ in their approach to time of 342flight measurements, and in the resulting instrumentation that this implies. TD systems 343emit extremely short pulses of light into tissue, and directly measure the arrival times of 344the scattered photons that emerge (Torricelli et al., 2014). Such recordings require very 345 sensitive photon-counting detectors. The time of flight multiplied by the speed of light in 346the tissue provides optical path length. In contrast, FD technique uses intensity-

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347 modulated light to illuminate the brain at very high frequencies, and measures both the 348attenuation and the phase delay of the emerging light (Wolf et al., 2007). Time of flight is 349then obtained by Fourier analysis of the phase delay, and can be used to calculate optical 350 path length. The resulting advantage of TD and FD imaging is that knowledge of optical 351 path length allows calculation of absolute values of HbO, HbR and total hemoglobin 352 concentrations. On the other hand, such systems are associated with higher costs, bulky 353 instrumentation, and slower acquisition times. The characteristics of the different fNIRS 354technique have been described in much greater detail in recent reviews (Wolf et al., 2007; 355Scholkmann et al., 2014; Torricelli et al., 2014).

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3573.2 Advantages, limitations and considerations for using fNIRS with CIs

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359 Compared to other techniques, fNIRS has several clear advantages that encourage its use 360in CI research. One of its most appealing features is its full compatibility with CI devices. 361 Owing to the optical nature of the technology, fNIRS data are not corrupted by the 362 electronic or ferromagnetic components of the CI device during acquisition. PET is the 363only other neuroimaging modality that provides a matching level of compatibility. 364However, unlike PET, fNIRS does not require injection of tracer substances in the blood 365 stream and does not expose individuals to radiation. The number of examinations is 366therefore not restricted, and repeat assessments through longitudinal studies can be 367 performed. fNIRS is also ideally suited for research involving young infants. 368Measurements can be recorded without the need for sedation or restraints because it is 369robust to motion artifacts. In fact, recording during overt speech in even possible (Hull et

370al., 2009; Quaresima et al., 2012). This is of great significance for CI investigators, as a 371large field of CI research involves the pediatric population.

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373Good research tools are safe, but also practical. To carry NIR light, fNIRS uses optic 374 fibers that are light, flexible, and therefore suitable for a range of head positions and 375 postures. Some centers replaced the plastic optic fibers with glass optic fibers and have 376 reported reduced weight of the optic bundles on the headgear (Lloyd-Fox et al., 2010). 377 Furthermore, fNIRS requires only a compact measurement system. The setup typically 378 consists of a mobile cart carrying a computer tower and monitor, an optical NIRS module 379 and the optical fibers connected to that module. This increases portability and allows for 380 measurements in non-intrusive environments and even in clinical settings. PET scans, on 381 the other hand, can only be performed in a radiation-proof radiological suite and require 382the presence of a radiochemist and a cyclotron for the production of radioisotopes 383(Crosson et al., 2010). Advances in optical technology have even allowed the production 384of a wireless, completely wearable, multi-channel fNIRS system suitable for use in 385 unrestrained settings (Piper et al., 2014). Cost is another important factor to consider 386when choosing a research instrument. fNIRS is among the most affordable neuroimaging 387 modalities, after EEG. There are no disposables and minimal maintenance is required. In 388 comparison, the instrumentation and maintenance fees associated with MRI, PET and 389MEG are on the order of millions of dollars (Bandettini, 2009).

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391 The temporal resolution of fNIRS is the highest among the hemodynamic neuroimaging 392techniques, reaching up to 100 Hertz (Hz) with CW systems (Huppert et al., 2006).

393Although inferior to EEG and MEG by one order of magnitude, this fine temporal 394 resolution allows the use of event-related paradigms and allows for nuanced examination 395 of the temporal dynamics of cortical blood flow. The spatial resolution of optical 396topography is typically estimated at 1 cm (Ferrari and Quaresima, 2012), enabling the 397localization of brain responses to specific cortical regions with reasonable precision. The 398 spatial resolution is dependent on the arrangement of source-detector fibers on the scalp. 399Increasing the density of channels, among other things, achieves finer sampling of the 400 cortex (Minagawa-Kawai et al., 2008). At our institution, we transitioned from a four 401 channel system to a 140 channel system, allowing us to generate topographic activation 402maps of the auditory cortex (Pollonini et al., 2014; Sevy et al., 2010). It is even possible 403to generate three-dimensional images of the optical properties of the brain given a 404 sufficient number of sources and detectors placed around the head (Minagawa-Kawai et 405al., 2008). This technique, called optical tomography, is costly and is usually restricted to 406 young infants, as adults' larger heads usually result in too much light attenuation (Gibson 407et al., 2005). Another advantage of fNIRS is that it offers quantitative monitoring of HbO, 408HbR, and total hemoglobin, generating a more complete evaluation of the cortical 409hemodynamic response than the fMRI BOLD response which tracks HbR (Scholkmann 410et al., 2014). Lastly, the fNIRS hardware is silent, which makes it ideal for the 411 presentation of accurate auditory stimuli in an acoustically-quiet environment, and 412 artifact-free response measurement.

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414The major spatial limitation of NIRS is that it only probes a thin top layer of the cortex, 415up to 1.5 cm deep (Fukui et al., 2003). This is a considerable drawback for cognitive

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416 studies that aim to investigate deep regions such as the brainstem, basal ganglia, or 417amygdala (Minagawa-Kawai et al., 2008). However, a substantial amount of research can 418be done probing the upper layers of the auditory, visual, somatosensory or frontal cortices 419 in CI research. Depth resolution is also highly dependent on the age of the subjects and 420 varies somewhat from region to region even within a particular age group (Beauchamp et 421al., 2011). In adults, thicker scalp soft tissues and skulls significantly restrict NIR light 422 penetration, impacting the accuracy of the recording. Deeper neural activity can be 423probed by increasing the source-detector distance, although at the cost of lower signal-to-424 noise ratio due to a reduction in the number of transmitted photons.

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426Good contact between the optodes and the skin of the scalp is also critical for a high 427 signal-to-noise ratio (SNR) and a good quality recording. Hair is a nuisance in fNIRS  $428$  recordings because  $(1)$  it interferes with this contact and  $(2)$  hair pigments significantly 429 scatter and absorb NIR light and therefore attenuate the detected signal. In subjects with 430thick, dark hair, a researcher can spend a considerable amount of time trying to optimize 431 the positions of the optodes to maximize the SNR. The use of gel can help to keep hair 432 pushed out of the way. Nevertheless, the best recordings often come from subjects who 433are bald or have thin, blond hair — this makes fNIRS particularly suitable for work with 434infants.

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436 Another drawback to fNIRS is the need to separate signals of cerebral origin from those 437 of extra-cerebral tissues. For instance, blood volume changes in the scalp and within the 438 muscles beneath the optical probes create noise in the fNIRS recordings and must be

439filtered during data analysis. Physiologic noise originating from heart rate and changes in 440 respiratory effort may also be a source of confounding cerebral blood flow signals and 441 must be accounted for during analysis (Gagnon et al., 2012). To remove the noise 442 component from the raw data, analytical strategies must be adopted. While some 443 institutions use their own custom software, others turn to freely available software 444 packages. However to date, there is a lack of a standard method for data analysis in 445fNIRS (Tak and Ye, 2014).

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447Similar to EEG, MEG and PET, the raw fNIRS data not provide an anatomic image upon 448 which neural activity can be superimposed. Therefore, to localize brain activity to known 449anatomical locations, the optodes must be carefully positioned according to a standard for 450the recordings. The 10-20 (EEG) system is often used (Minagawa-Kawai et al., 2008). 451 Once this is done, the optode layout is precisely aligned, and therefore the functional data 452 obtained with fNIRS can be overlaid onto structural MRI images or anatomical atlases, if 453 desired (Crosson et al., 2010).

#### 454

455 Certain considerations must be taken into account when acquiring fNIRS data from CI 456 users. Depending on the probe layout and the size of the headset, the external magnet of 457 the CI device can interfere with headset placement over the temporal area. In such 458 circumstances, we simply place the headset over the magnet (Figure 1). While this 459 obstructs the scalp contact of certain channels, the remaining channels can still be used. 460In our experience, however, the external magnet is generally posterior and inferior 461enough so as not to interfere with headset placement that permits the measurement of

462 responses within the regions of interest, such as primary auditory cortex. Of course, care 463 must be taken not to displace the magnet, as the implant would turn off. Gentle 464 manipulation is also required when placing the headset in the crease between the pinna 465 and the temporal skin to avoid repeated contact with the CI microphone and the resultant 466unpleasant noise for the CI user.

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468In an attempt to facilitate recording in the CI population, we designed a custom probe 469layout and headset at our institution. This arrangement features six light sources clustered 470in the center of the headpiece and an additional source anteriorly and posteriorly. 471Detectors are positioned in between (Figure 2D). The center-to-center distance between 472adjacent optodes was 15 mm. Moving away from the checkerboard pattern described in 473our previous work (Figure 2C; Pollonini et al., 2014), this new honeycomb-shaped design 474 allows for a denser configuration of probes, while maintaining an equal number of 475 channels. The result is a smaller and more convenient headpiece suitable for both adult 476 and pediatric subjects, without compromising resolution. This dense multi-array headset 477 allows spatial oversampling of a defined cortical area through adjacent channels that 478 cross each other.

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4803.3 What region(s) of the central nervous system should be studied?

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482To understand the neural substrates involved in auditory processing through cochlear 483 implants, it is necessary to observe activity within the brain when a sound stimulus is 484 presented (Hall and Langers, 2014; Zhang et al., 2010). Ideally, one would track activity

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485all the way from the level of the auditory nerve, through the ascending auditory pathways 486in the brainstem to the auditory and auditory-associated cortical regions. However, given 487its depth limitations, such whole-brain imaging is not possible with fNIRS. Because 488fNIRS is not a whole-brain technique, choices must be made about what portion of the 489 cortex to record from in order to get the information most relevant to understanding 490 auditory processing through a CI. A substantial body of fMRI data highlights the lateral 491temporal lobe and superior temporal gyrus (LTL/STG) as foundational to auditory 492 processing at the cortical level.

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494 Several studies have revealed preferential activity for the processing of acoustic 495 parameters such as pitch, noise and spatiotemporal fluctuations in the LTL/STG (Hall and 496Plack, 2009; Humphries et al., 2010). Selective responses to species-specific 497 vocalizations were demonstrated in the LTL/STG of humans and other mammals (Belin 498et al., 2002). In addition, studies using fMRI and implanted recording electrodes have 499shown localized responses within the left LTL/STG to phonemes, words, and phrases 500(DeWitt and Rauschecker, 2012). Of particular relevance to understanding hearing 501through a CI, Smalt et al. (Smalt et al., 2013) demonstrated rapid neural adaptations in 502normal-hearing participants exposed to degraded sound, similar to what a CI user 503experiences.

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505 While fNIRS does not provide whole-brain imaging, it can be used to dissociate music 506 and language processing within constrained cortical regions such as the left and right 507LTL/STG thanks to stimulus specific processing differences across the cerebral

508hemispheres. Neuroimaging studies in normal-hearing subjects using PET and fMRI have 509 previously shown that the left temporal lobe is primarily involved in speech and language 510 processing, while the right temporal lobe preferentially responds to music (Hickok and 511Poeppel, 2007; Price, 2000; Belin et al., 1998). Furthermore, reports have demonstrated 512that secondary auditory areas in the right STG (surrounding Heschl's gyrus) are key to 513the processing of pitch information (Zatorre, 1998; Tramo et al., 2002). Temporal 514information, on the other hand, is preferentially processed by left-lateralized primary 515(core) auditory areas (Zatorre and Belin, 2001). Evidence also points toward a functional 516 segregation between music and speech processing within the temporal lobes (Abrams et 517al., 2011; Levitin and Menon, 2003). Armony and colleagues not only revealed the 518 existence of a region in the anterior STG (planum polare) that responds more strongly to 519 music than voice, but their results also provide strong support for the presence of "music-520preferring" neurons in this area (Armony et al., 2015). Moreover, several fMRI studies 521 have demonstrated that the anterior portion of the STG is involved in higher-order music 522 analyses such as extraction of melodic information (Rogalsky et al., 2011). Lesion studies 523 have reinforced the idea that pitch and rhythm processing recruit separate neural 524 subsystems within the auditory cortex: cortical damage can interfere with pitch 525 discrimination without affecting rhythm performance, and vice-versa (Di Pietro et al., 5262004; Ayotte et al., 2000). These and other findings indicate that the LTL/STG are the 527 most clinically relevant regions of the cortex to focus on when imaging different classes 528 of auditory perception in CI recipients using fNIRS.

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5303.4 Data analysis techniques in multi-array fNIRS headsets

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532A comprehensive review of analysis techniques available for use with fNIRS data is 533beyond the scope of this paper, and this topic has been extensively reviewed recently 534(Tak and Ye, 2014). Rather, in the following section we summarize current strategies to 535 analyze recordings from dense multi-array headsets, as they are the most suitable for CI 536 research. As with fMRI, signal pre-processing is initially performed to remove motion 537 artifacts and physiologic noise. The first step requires identification of channels with 538good scalp contact. At our institution, we filter channels with excessive noise according 539to their scalp-coupling index (Pollonini et al., 2014). In brief, this technique relies on the 540fact that adequate scalp contact is characterized by a synchronous cardiac pulse signal 541 recorded by both wavelengths of light emitted from a single probe. While a perfect 542 correlation between each wavelength's cardiac signals is ideal (coefficient of 1), channels 543 with an index threshold above 0.70 are reliable and can be retained.

#### 544

545The next step is motion artifact correction. Relative to hemodynamic-related changes, 546head movements will cause rapid changes, sharp spikes, and increases in the magnitude 547 of the recorded signals (Tak and Ye, 2014). Previous reports have described the use of 548 external accelerometers to estimate and correct baseline motion artifacts, but this requires 549 additional instrumentation with its related cost and complexity (Virtanen et al., 2011). 550Many approaches to remove these artifacts without the need for motion sensors have also 551been described (Cui et al., 2010; Scholkmann et al., 2010). Our preferred technique 552 consists of identifying start and stop times of motion artifacts by bandpass filtering each 553 channel between 0.1-3.0 Hz to remove slow signal drift and by normalizing the intensity

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554 of the highest peak of the entire time course. We define peaks in the signal exceeding 55520% of the maximum peak intensity as motion artifacts. These are then removed from the 556 raw data by performing linear interpolation between the start and stop time points. Once 557 motion artifacts are corrected, physiologic noise can be removed from the hemodynamic 558 signal. This is usually accomplished by bandpass filtering between 0.016-0.25 Hz. The 559 modified Beer-Lambert law is then used to calculate the relative concentrations of HbO 560 and HbR for each channel and time point (see Section 3.1).

#### 561

562 Once signal processing is complete, brain activation can be detected by performing 563 inferential statistics on the fNIRS data. For each channel, all the trials of each stimulus 564 first need to be averaged, a process called block-averaging (Scholkmann et al., 2014). 565The resulting block-averaged hemodynamic response is then compared to a predicted 566hemodynamic response. Predicted fNIRS responses can be modeled in a manner similar 567to the analysis of fMRI data (Cox, 1996). In such models, the HbO concentration rapidly 568rises after stimulus exposure, reaching a peak in a few seconds. The response then 569 plateaus pending stimulus discontinuation, following which it slopes down until baseline 570HbO concentration is reached. Physiologically, this corresponds to an augmented blood 571 supply required by the neuronal activation. Conversely, HbR concentration changes in a 572 similar but opposite direction, decreasing during stimulus presentation. The quality of fit 573 is determined by linear regression analysis of the measured and predicted 574 responses, resulting in a T-statistic for each channel. Thus, each source-detector pair 575(channel) in the headset can be represented by a single number that describes the 576 goodness of the fit. Thes T-statistics are then arranged in a spatial grid representing the

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577 position of the channel they derive from within the source-detector array. Multi-array 578fNIRS headsets provide spatial oversampling in the cortex since many channels cross 579 each other at a given location. The resulting benefit is a reduction of noise in overlapping 580 channels. A topographic (2 dimensional) activation map for each stimulus condition can 581 then be generated by color-coding the T-statistic spatial grid. Alternatively, it is possible 582to project this colored T-statistic distribution map onto a standard brain image to create 583 cortical activation maps that are easier to visualize and interpret.

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#### **4. Review of fNIRS neuroimaging studies in CI recipients** 585

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 $587$ In 2013, fNIRS celebrated its  $20<sup>th</sup>$  anniversary as a human neuroimaging modality. Jöbsis 588(1977) was the first to demonstrate the possibility of detecting changes of cortical 589oxygenation by transilluminating the cranium of anesthetized cats with NIR light (Jöbsis, 5901977). However, it was not until 1993 that this emerging technology was first applied to 591human brains. That year, four research groups independently published the first single-592site fNIRS human adult studies (Chance et al., 1993; Hoshi and Tamura, 1993; Kato et 593al., 1993; Villringer et al., 1993). fNIRS has since rapidly gained popularity among the 594 neuroscience and clinical communities. If the number of annual publications reflects 595 scientific enthusiasm, fNIRS has definitely emerged as one of the most popular research 596 fields in the past 20 years: its publications have doubled every 3.5 years and have now 597 reached over 200 per year (Boas et al., 2014). Despite this growing interest, the literature 598 reporting the use of fNIRS in the CI population remains sparse. A comprehensive review 599 across multiple databases of published articles mentioning fNIRS and cochlear

600implantation yielded four papers (Sevy et al., 2010; Pollonini et al., 2014; Dewey and 601Hartley, 2015; Lawler et al., 2015) and one conference abstract (Olds et al., in press). 602

603Sevy and colleagues report the first research application of fNIRS in CI users (Sevy et al., 6042010). The authors used fNIRS to measure speech-evoked cortical responses within four 605 subject cohorts: normal-hearing adults, normal-hearing children, deaf children who had 606 over 4 months experience hearing through a cochlear implant, and deaf children who 607 were tested on the day of initial CI activation. The speech stimuli consisted of digital 608 recordings from children's stories in English. A four channel NIRS 2CE system (TechEn, 609Inc., Milford, MA) with 2 emitters mounted on a custom headframe was used to sample 610bilateral auditory cortices (Figure 2A). The authors report successfully recording auditory 611 cortical activity using this fNIRS setup in 100% of normal-hearing adults, 82% of 612normal-hearing children, 78% of deaf children who have used a CI for at least four 613 months and 78% of deaf children on the day of CI initial activation. Interestingly, Sevy et 614al. had validated their NIRS experimental paradigm with fMRI in 3 normal-hearing 615 adults. They showed that similar speech-evoked superior temporal gyrus responses were 616 obtained with both fNIRS and fMRI. Such results were encouraging as they demonstrated 617that fNIRS was a feasible neuroimaging technique in CI users and that reliable 618hemodynamic cortical responses to speech could be recorded in these patients.

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620The same group later evaluated whether fNIRS was sensitive enough to detect differences 621in cortical activation evoked by different quality levels of speech in normal-hearing 622individuals (Pollonini et al., 2014). The investigators used a 140 channel fNIRS system

623(NIRScout, NIRx Medical Technologies LLC, Glen Head, NY) in a tight array to provide 624 spatial oversampling, and permit averaging between channels to improve the SNR 625(Figure 2C). By increasing the number of channels, the authors were able to generate 626topographic maps and measure the area of activation and center of mass. They also 627 designed their own custom analytic software and developed novel data analysis 628techniques to filter channels with poor scalp contact or high SNR. The experimental 629 paradigm consisted of four different stimuli: normal speech, channelized (vocoded) 630 speech, scrambled speech and environmental noise (for previous use of these stimuli as 631 cross-controls see, for example, Abrams et al., 2011; Humphries et al., 2001; Levitin et 632al., 2003). Their results revealed that speech intelligibility correlated with the pattern of 633 auditory cortical activation measured with fNIRS: normal speech evoked the strongest 634 responses, distorted speech produced less region-specific activation and environmental 635 sounds evoked the least response. Again, the investigators validated their stimulus 636 paradigm with fMRI on a single participant. Such results demonstrated that in normal-637hearing individuals, fNIRS can detect differences in the response of the auditory cortex to 638 variations in speech intelligibility. The conclusions of this study raise implications for the 639CI population. If fNIRS can provide an objective measure of whether a normal-hearing 640 subject is hearing normal or distorted speech, then it has the potential to be used to assess 641 how well speech information activates the brain in subjects hearing through a CI.

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While Pollonini's study did not involve CI subjects, subjects hearing through a CI were 643 644 studied with a similar technique (Olds et al., in press). Olds' study used an experimental 645 paradigm and fNIRS instrumentation comparable to that of Pollonini, but expanded the

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646 approach to participants with CI. Specifically, the authors aimed to better understand the 647 variability in speech perception outcomes in CI using fNIRS. A NIRScout 1624 648instrument (NIRx Medical Technologies, LLC, Glen Head, NY) with 140 channels was 649used to record the auditory cortical response of 32 post-lingually deaf adults hearing 650through a CI and 35 normal-hearing adults. Again, four auditory stimuli with varying 651 degrees of speech intelligibility were employed: normal speech, channelized speech, 652 scrambled speech and environmental noise. Speech reception thresholds (SRT), 653 monosyllabic consonant-nucleus-consonant word (CNC Words) scores and AzBio 654 sentence recognition scores were used as behavioral measures of speech perception. 655 Results from this study demonstrated that the cortical activation pattern in implanted 656 adults with good speech perception was similar to that of controls. In those two groups, 657less cortical activation was noted as the speech stimuli became less intelligible. In 658 contrast, CI users with poor speech perception displayed large, indistinguishable cortical 659 activations across all four stimuli. As the authors had hypothesized, the findings of this 660study demonstrated that activation patterns in the auditory cortex of CI recipients 661 correlate with the quality of speech perception. Importantly, when the fNIRS 662 measurements were repeated with the implant turned off, reduced cortical activations in 663all CI recipients were noted. This suggests that sound information is conveyed to the 664 auditory cortex of CI users with poor speech perception, but that these subjects are unable 665to discriminate speech from the information that gets to the cortex.

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667To our knowledge, Lawler and colleagues are the only other research group actively 668using fNIRS neuroimaging in auditory processing studies in deaf individuals and CI

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669 recipients; to date, they have published two articles on that topic (Dewey and Hartley, 6702015; Lawler et al., 2015). While this group's long-term aim is to examine cortical 671 reorganization associated with deafness and cochlear implantation using fNIRS, none of 672 these articles enrolled CI users thus far. The first report discusses maladaptive cross-673 modal plasticity in CI subjects and its role as a potential factor underlying poor 674 performance following implantation (Lawler et al., 2015). Through this article, the 675 authors describe their long-term research goals and introduce their plans for future fNIRS 676 studies with deaf individuals and CI recipients. Later that year, Dewey and Hartley 677 published a study on the use of fNIRS to detect visual and vibrotactile cross-modal 678 plasticity changes in profoundly deaf but non-implanted individuals (Dewey and Hartley, 6792015). Their setup consisted of a Hitachi ETG4000 (Hitachi Medical Corporation, Tokyo, 680Japan) optical topography system with 12 recording channels over each hemisphere 681(Figure 2B). The authors reported that auditory deprivation is associated with cross-682 modal plasticity of visual inputs to auditory cortex. Practically speaking, such results 683highlight the ability of fNIRS to accurately record cortical changes associated with neural 684 plasticity in profoundly deaf individuals. The application of these findings to the CI 685 population is very promising, as they demonstrate the potential of fNIRS as an objective 686 neuroimaging tool to detect and monitor cross-modal plasticity both prior to and 687following cochlear implantation.

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#### **5. Directions for future fNIRS application in CI users**  689

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5.1 Clinical applications 691

693A promising future for fNIRS clinical applications includes the implementation of NIRS 694as a neuroimaging tool to guide post-implant programming in the service of improving 695 deaf patients' speech and language outcomes. CIs need to be reprogrammed frequently to 696 ensure they are accurately conveying the sound information within speech to the auditory 697 nerve and, ultimately, to the auditory cortex. If the language areas of the brain are 698 appropriately activated, then the child has the best chance of learning normal speech and 699language. Early identification of patients who do poorly is therefore critical, as prompt 700intervention can prevent delay in linguistic and psychosocial development (Robinshaw, 7011995). Current cochlear implant assessment tools are limited and hard to administer in 702young infants, whose behavioral responses are difficult to elicit and are often not 703 interpretable. An objective measure of how well speech information is processed within 704the cortex would provide an ideal tool for monitoring (and possibly predicting) language 705 development in young CI users. Given that the number of imaging sessions is not 706 restricted for fNIRS, repeat assessments through longitudinal studies can be performed to 707 monitor rapid cortical modifications resulting from poor implant programming. In doing 708so, fNIRS studies may allow early identification of children on poor language 709 development trajectories. If this can be achieved while the child is still within the critical 710time period when significant language development occurs (i.e. age 1-4 years), prompt 711 intervention can be started. Ultimately, this type of early intervention could prevent 712 delays in a child's psychosocial development, a process highly dependent on hearing 713(Yoshinaga-Itano et al., 1998). Using fNIRS to supplement our current clinical practice of 714CI programming and speech and language therapy is an exciting possibility.

#### 7165.2 Research applications

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718The opportunity for safe, repeated testing of CI recipients with fNIRS also provides 719 investigators with the ability to explore the cortical changes associated with neural 720 plasticity in this patient population. For instance, understanding the cortical 721 reorganization that occurs following prolonged auditory deprivation in potential CI 722 recipients may help predict their expected outcome post-implantation. This expectation is 723based on emerging evidence suggesting that cross-modal plasticity of visual inputs into a 724 sensory-deprived auditory cortex may affect the ability of a CI recipient to process 725 auditory information from their implant effectively (Sandmann et al., 2012). fNIRS may 726also provide insight into the cortical changes that take place in deaf patients following 727implantation. An example of such an application is the study of post-implantation training 728 and its effects on brain plasticity. Pantev et al. examined the dynamics of auditory 729 plasticity after implantation through MEG longitudinal imaging, suggesting that CI users 730 would benefit the most from language training within the first 6 months after 731 implantation (Pantev et al., 2006). As discussed, fNIRS is significantly easier to use in 732longitudinal studies compared to MEG. The opportunity to further explore cortical 733 reorganization following hearing restoration has the potential to guide the design of post-734 implantation training strategies.

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736 The neural basis for CI users' variable experience perceiving music is another interesting 737topic and one that merits further investigation. Despite advances in CI technology, music

738 perception in CI recipients remains quite poor (Gfeller and Lansing, 1992). A growing 739body of psychophysical studies has better defined the limitations of music enjoyment and 740 perception in CI users. For example, studies suggest that CI users perform poorly on pitch 741 recognition tasks, whereas rhythmic perception remains relatively intact following 742implantation (W. B. Cooper et al., 2008; McDermott, 2004). Reports have also shown 743that appraisal ratings and overall listening time are significantly lower following 744implantation, with some CI users even describing music as "aversive" (Looi et al., 2012; 745Migirov et al., 2013). The challenges that CI users face in processing a complex auditory 746 stimulus such as music can be explained by a number of technological, acoustical and 747biological constraints (Limb and Roy, 2014). While many of these have been addressed in 748the literature previously, the neural basis for poor music perception in CI users is under-749 investigated and poorly understood. This is at least in part due to inherent limitations on 750the use of most neuroimaging modalities with CI users, as outlined here. fNIRS is quiet 751 and allows the use of event-related paradigms, thus offering greater flexibility in 752 experimental inquiry. It is also relatively low cost, another factor that may have 753 constrained examination of neural mechanisms underlying better or worse music 754 perception in implant users in previous years. These and other features make fNIRS an 755ideal tool for evaluating music-evoked brain activation in CI recipients, as well as for 756 examining the relationship between behavioral music performance and degree of auditory 757 cortical activation in this patient population. Together, these inquiries would help achieve 758the long-term goal of higher-level music perception in CI recipients.

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7606. Conclusion

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762fNIRS is a safe, reliable neuroimaging technique that is compatible with CI devices. It

763 offers many benefits over other approaches for examining cortical responses in CI

764 recipients, although care must be taken in collecting and analyzing the data. While the

765 existing literature on fNIRS neuroimaging in adult and pediatric CI users is currently

766limited, the future of this emerging technique is promising and numerous clinical and

767 research applications remain to be explored.

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fNIRS: functional near-infrared spectroscopy, fMRI: functional magnetic resonance imaging, PET: position 1306 emission tomography, EEG: electroencephalography, MEG: magnetoencephalography 1307 Figure 1. fNIRS headset placement over a cochlear implant device. A) The location 1304 1305

1308 of the cochlear implant's external magnet and coil interferes with headset placement over

1309the temporal area. B) The fNIRS headset is simply apposed over the magnet (shaded 1310area). C) Diagrammatic representation depicting the quality of scalp contact of the optode 1311 array, obtained from custom analytic software using real-time fNIRS recordings. The 1312 optodes obstructed by the magnet postero-superiorly lose their scalp contact (red), while 1313the remaining optodes are unaffected and can still be used (green). The status of scalp 1314 contact was indeterminate for certain optodes (yellow).

### **Figure 2. Comparison between fNIRS probe layouts previously reported for CI use.**

1346A, Sevy (2010); B, Dewey (2015); C, Pollonini (2014); D, Our new honeycomb-shaped

1347headpiece. The optode arrangement in all headsets is based on the International 10/20

1348 system: A is centered at the T3/T4 position; the optode located in the middle of the

1349 bottom horizontal line in B, C and D is aligned with the T3/T4 position.