A Letter to the Professional Community

Dear Professional,

Since my daughter was diagnosed as being profoundly deaf over five years ago, I have sought and strived to gather together medically proven and factual information on how to best teach a deaf child to hear and speak very, very well. In our case, there were no clear answers, and too many opinions prevailed. Having about two weeks of education under my belt, I was asked to choose a language modality to best fit our family and my deaf child. There was no detailed information of the varying language modalities and certainly no idea of what the end result could be with whatever option we chose. All I knew was that oral speech was the goal for my deaf child, but precisely what needed to be done did not exist. In this day and age five years later, I am certain it still does not exist. This is why this website and the attached PDF files are here.

So with this said, within the text of the PDF files are very, very clear road maps of everything that needs to be in place and everything that needs to be done for an extremely successful outcome for parents and professionals who choose a 100% oral option for their young deaf child. This book showcases step by step exactly what a parent needs to do, and the programs and professionals that need to be in place so a profoundly deaf child can obtain “Perfect Speech, Audition, and Language.” I realize that this is a profound medical statement, but let’s not kid ourselves. In this day and age deaf children CAN achieve absolutely normal language with no deficits at all and in many cases with much higher cognitive and language abilities than hearing children. (This may not be possible with all children due to certain medical conditions or lack of needed services.)

You may ask yourself how can any guidelines apply to every deaf or hard of hearing child? Just like every car needs the oil changed, every car needs the tires changed, and every car needs frequent tune-ups, in my mind every child who is deaf / HOH can benefit from very specific guidelines and procedures.

The real question is how is this done? How do we get from point A to point B to obtain the best possible medical outcome? There is nothing I can find on the internet or in any case study which explains “What to do.” Many pamphlets and organizations will tell you the best method or program, but then what? To lay out EXACTLY what to do for the next three to five years until that child hears and speaks identically to a hearing child does not exist. In my book it does. Yes, every child is different and there is no guarantee. I have heard it a hundred times. If you look at the line items and read Chapters 2, 4, 6, and 7 and apply EVERYTHING that is mentioned, no deaf child could fail, and only the absolute best medical results will be achieved. Will every child have “Perfect Speech and Language?” Maybe, or maybe not, but the ABSOLUTE best results possible for that child will be achieved. This is the question the medical community has been trying to answer with study after study after study. This is answered in the book. These chapters in my opinion need to be studied for their medical merit. Not one professional, organization, or book ever explained to our family what is written in these four chapters. Keep in mind that this book is only written for parents who have already discussed with their doctor / SLP and have chosen oral language as the appropriate modality for their child.

You are on the medical / professional side with excellent resources and study material. I am on the parent side with the experiential equivalent of a four year degree in “How to Teach a Deaf Child to Hear and Speak.” This degree has no price tag, but is formed and earned with motivation, desire, and need. There are two sides to this equation which hold equal weight: the professional side and the parent side. For the best results with any child both entities must be in complete harmony.

There are many fine professionals and many dedicated parents who put everything they have into the habilitation and lives of their deaf / HOH children. The reason in my mind to lay out clearly what to do is to help the young deaf / HOH child achieve excellent language, audition, and articulation so they can reach for
their goals, which will form in the child’s teenage and young adult years. It hurts me to see any child with poor articulation, language, or cognitive skills when the end result should have been different.

The problems as I see them and the reasons many children do not reach the highest possible and best outcome with oral education are as follows:

1) Parents are not educated sufficiently and / or motivated. The key is to make the RIGHT choices without making mistakes. Yes, parents will make a choice but is it the RIGHT choice?
2) Parents do not make the needed sacrifices as laid out in Chapter 2.
3) Parents do not seek out and receive early services or implantation.
4) Parents listen to one opinion and take action without considering all options.
5) Parents do not do what is absolutely necessary at home to fully help their deaf child hear and speak.
6) Parents do not have the tools, books, or programs available to them as laid out in Chapter 9.
7) Professionals do not have the motivation to hand parents a 3 – 5 year guide / checklist such as I have written.
8) Wrong medical advice still exists, such as:

   a) Suggesting Total Communication or ASL for early implantees with no other medical conditions.

   b) Making the statement that “If the child is making good progress with the hearing aids then leave them on.” The real question should be; will the child make better progress at 15 dB with a cochlear implant or 45 dB with a hearing aid? This in my mind is self-explanatory so why is the above statement still made?

   c) Professionals providing or going along with inadequate services without making the parent knowledgeable enough to seek out better opportunities for their child.

There is NO absolute, clear road map for a parent to follow. It is left to parents to find their own way. Once a parent makes the decision for oral language, where are the written, step-by-step instructions which can be used for the next three to five years?

Opinions from professionals are just that, opinions. In the very beginning I would ask three people what to do and they all had their own opinion. I do not favor opinions. Since my daughter was diagnosed I wanted to know what to do from previous medical research. Tens of thousand of children have obtained oral language from numerous methods. I would ask; “what is the best method for my daughter to obtain oral language?” The answer would be: “Well Mr. Hall, you need to make the decision of which language option will work best for your family.” Not the answer I wanted. The answer I wanted was: “Of the tens of thousands of children who have obtained oral language in the past five years we studied language abilities, audition abilities, articulation, discrimination, comprehension, cognitive skills, and reasoning, and we have found that of these tens of thousands of children the highest ranking individuals used the XYZ method.” 100% factual with no opinions. It does not exist, I never received any statement close to this, and why it does not exist in the medical community I do not understand.

In closing, I have attached the following list of procedures for your reference. I am trying to present this list to the medical and educational community as a guideline for parents and professionals who have chosen the 100% oral-only option for their young implanted child. This list is not extensive, it is only seven pages, the full text (48 pgs) from which this list was drawn is in Chapters 2, 4, 6, and 7 of A Father’s Love, provided on this website FREE OF CHARGE. A three-page summary of the book can also be found on this website under the link “How to Use This Parents’ Manual.”
It is frustrating to me that hundreds and even thousands of parents and professionals are seeking the same answers over and over again and making many mistakes along the way, when there is a clear road map, it only needs to be followed. Please read the following seven-page list and even Chapters 2, 4, 6, and 7. Make changes or disagree, it may or may not be perfect, but this is the only clear step-by-step procedure of what to do over the next three to five years that I have found.

I am asking for your help to please reply to this letter at the attached email address to see why the following list and Chapters 2, 4, 6 and 7 cannot be presented as a medical guide for young cochlear implanted patients / students.

Please keep in mind that what is written is not opinion. Many deaf children can reach age-appropriate language by year two or three post-implant. What is written is 100% factual of everything that was done so our deaf child could:

Obtain above age level language.
Obtain above age level cognitive skills.
Become fully mainstreamed with no supportive services from kindergarten forward.
Answer any open set question in complete darkness.
Respond appropriately to a three-part question with no visual cues.
Respond to a whisper and answer appropriately to any open set question with no facial cues.
Discriminate 5 – 10 rhyming words with 100% accuracy auditory-only.
Read written words and sentences at age 3½.
Know her entire phonetics by age 3½.
Be in the 97th percentile for cognitive and language skills.
Have completely normal clear language and articulation with absolutely no deficits.

This is not an isolated case. My goal is to see more children reach such milestones.

Thank you in advance for your response.

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How to Obtain the Absolute Best Medical Outcomes for Children Using Oral Education

The goal of everything I write in these pages and at the website www.deafchildrencanspeak.com is to help parents and professionals obtain the “absolute best medical outcome” for every deaf / HOH child. There are certain things that apply to every child going through the oral process. When all of these items are maximized to their full potential it becomes an extremely powerful habilitation plan to obtain the best results.

Just like every car needs the oil changed, every car needs the tires changed and every car needs frequent tune-ups, in my mind every child who is deaf / HOH can benefit from very specific guidelines and procedures.

The following are excerpts from Chapters 2, 4, and 7 of A Father’s Love, which can be downloaded FREE OF CHARGE at the above website.

I am a father of a profoundly deaf child who in my opinion has obtained the absolute best medical outcome possible by following the guidelines and procedures listed below. This young girl can follow any open set question with no visual cues, is above age level in language, has perfect articulation, and has the ability to blend in with the hearing world with absolutely no one having any idea she is profoundly deaf. I am hoping for the best results for your child. In my opinion, that means the correct procedures must be followed from the very beginning. This is why most of my efforts are concentrated on the very young population of deaf / HOH children aged 24 months and younger. Do keep in mind that these guidelines will be beneficial for the next three to five years as you go through this language and audition journey with your young child.

The medical community has been trying to answer the question of “the best method” for years. Study after study after study is done to find out “the best way to teach deaf children.” If criteria such as audition, articulation, language abilities, hearing thresholds, and other items are taken into consideration, the procedures below are the answer to “the best method.” For families who choose oral speech, these methods only need to be followed.

The following list has worked wonders in our family. It may or may not be complete. But please look at this list and ask yourself what would happen if every piece was in place to help every deaf / HOH child? This list is extremely powerful when maximized to its full potential. Above all put your entire heart and life into this process for the next three to five years and excellent results will follow.

1) For severe to profoundly deaf children, early cochlear implantation between the ages of eight and fifteen months of age. You need to have a very well qualified cochlear implant surgeon and pediatric audiologist on your team. This also raises the point of early diagnosis. If you even think your child is having a problem hearing please have a hearing test done earlier and not later.

2) A parent driven to give their child everything needed to succeed.

3) A parent who spends every waking hour learning more and more about language development and the oral habilitation process. This would include going to numerous conferences, reading numerous books, attending the speech therapy sessions, and talking to many parents and professionals.

4) A good auditory oral school or a program that resembles what the school would provide. Enroll your child as early as possible, preferably before twelve months of age. Most schools and Early Start programs will want to start parent education and formulate a program for a child as early as possible.
5) A school district /Early Start program which adheres to the IDEA law and can see that deaf children can learn normal speech and language. A district willing to fund four or more years of education. This could be as much as $50,000 (based on 2006 dollars). A district willing to work with you and understand the vision that you have for your child. In a lot of circumstances all it takes is to let people see the success of other children who have gone down the oral path, and it may change a school district’s / Early Start administrators’ mind in regard to funding a particular program.

6) Never take one professional’s opinion. Find out for yourself by asking numerous questions of numerous professionals and decide on your own gut feeling what is correct. (This author is available to help weigh the pros and cons of any of the numerous life-changing decisions in this process.)

7) Purchase the Ausplan book or a similar language development book and be in sync with the speech therapist. Reinforce all therapy in a natural setting at home. Know where your child is in acquiring normal language and know what the next step is. This ability to be able to track your child’s progress and pinpoint where they are on the language hierarchy model is so crucially important. (If you do not know this you cannot spend the numerous hours needed at home to reinforce what is being taught in the speech therapy sessions.)

8) About two to three years into this process provide a mixture of oral training and mainstream classes. Provide your child with age appropriate language models including fully hearing children to converse with.

9) Get your deaf child around as many hearing children as possible. Enroll your child in numerous preschool classes to help with their auditory skills, such as music, art, playgroups, etc. Their language and social skills will be enhanced by being around other children. Young normal hearing children are such great language and speech models. Some auditory oral programs even make it a point to have a full-time hearing student in the classroom to help model and facilitate language.

10) Be thankful, pleasant, and truly appreciative of everything every professional is doing for your child. Buy gifts, give cards, and let these people know they are helping change your child’s life for the better. These talented individuals are giving your son or daughter the gift of speech that no one can ever take away. Show your appreciation.

11) Have a family support network of at least one grandparent, brother, sister, uncle, or aunt who is dedicated to this process for the four years it will take. This means driving your child to school, the therapists, the doctors, and babysitting as needed. For a single parent or a married couple to put in the time and effort needed in this process is very difficult. This is why one additional truly dedicated family member is so very important.

12) Do your homework and take every IFSP / IEP meeting seriously. Know the IDEA law and put a positive spin on why the local school district / Early Start program should fund three to four years of oral education for your deaf child. It is so vitally important to consult with professionals who have gone through numerous IFSP / IEP meetings and understand the IDEA law. Using the right terminology and constructing your arguments in a manner that is backed up by the IDEA is so very important. Make the process cost-effective for the school district.

13) Flextime for one parent. To have a job which gives you the ability to go to the speech therapy sessions, the doctor appointments, the IFSP / IEP meetings and truly be involved in your child’s habilitation plan. Trying to go through this process with a boss counting every single minute you need to take off to be with your child just does not work well. You need a boss who wants to help you help your child and give you a flexible schedule. The exact verbiage of how to ask for flextime is listed in Chapter 2 of “A Father’s Love.”
You as the parent are in the driver’s seat. You must make the right decisions at the right time for the absolute best outcome for your child. This may seem odd, with numerous professionals who have years and years of experience dealing with deaf children. But in the beginning you will be a baby needing to be fed information. As time goes on you must be a lion who knows what they want and goes after it. You are the quarterback with a team of professionals all working toward the best outcome for your child. It is your responsibility to make sure that every professional is in sync with needed information and that these people talk to each other and the team work seamlessly for your deaf / HOH child.

**Things the Parents Need to Do at Home**

The following is what is needed in the beginning of the oral process to help your child speak. You will learn what needs to take place at home to give your deaf child the best possible outcome. Like you, I wanted my daughter to order her own food, ask for directions, converse on the phone, and have a “normal” social life. I did not want her to be secluded to the 1% of society who knows ASL. I wanted her to converse articulately with the 99% of society who use spoken language.

The very first thing that you need is a good quality oral program for your child. Do not let the school district / Early Start program decide for you. You have the right to a private oral school which the school district will pay for if they do not have an “appropriate” program in place. You can locate most of the oral schools in the U.S. at [www.oraldeafed.org](http://www.oraldeafed.org) Call this organization and request some of their excellent videotapes / DVDs, which have amazing cochlear implanted children who speak nearly perfectly. If you are not close to a school, find a good auditory oral / verbal therapist in your area. In addition to going to an auditory oral school and therapy, you need to do the following at home every day with your deaf child to facilitate speech.

1) Point out every sound, i.e., the phone rings and you say “I hear that, that is the phone” and point to your ear. If she is eating, say “You are eating apples,” etc. Everything, all the time, narrate what your child is doing and what they are playing with. “You are playing with Elmo, Elmo is red.” Narrate what you are doing. “Look, Mommy is cooking. I take the pan and put it on the stove…” Repeat the words dozens and dozens of times throughout the day. Whether it be the phone, door, birds, car, vacuum cleaner. Whatever the sound is, point to your ear and say “I hear that, that is the _____.”

Keep in mind that children need to learn to listen before they can speak. This goes back to proper therapy. A lot of time needs to be spent on listening before they can graduate to spoken language.

2) Cut out ALL background noise as much as possible. I love music and used to listen to music every day. Over the past four years I have drastically reduced how much the stereo is on because of the competing background noise. You need to make a few sacrifices in this area and divide the day into two distinctive parts: the time you work with your child, and the time you watch TV, listen to music, vacuum, etc. If the family has the TV on 24 / 7 in the background you must make a sacrifice in this area so you can have quality “quiet” time with your child. Have the washing machine and dishwasher going when you are away from home or the same time you watch TV. Make everything a quality listening experience.

3) If there are other children in your family have these children play with your deaf child as much as possible. These hearing children will provide an age-appropriate language model that your child can learn from. Let them play games, talk about games, talk about the weather, fight, yell, and do all the things young children do. Don’t sit there and have all your children glued to the TV set. The more quality speaking your child gets from siblings will really benefit their language development. Television is good if used properly to play age-appropriate language-rich DVDs and videotapes.

It is very important that our children be in play groups and other social events with hearing children. Try to have your child around hearing children as much as possible. This will help with
their speech and language.

4) Speak to your child all the time. Again, look directly at their face so they can see your mouth. Explain everything to them. Be within 3 - 4 feet when speaking and just talk, talk, talk, in English. This is what these kids need—a huge input of quality language so they can record everything in their auditory memory. Your therapist will be working on auditory-only exercises and you will soon be able to speak with your young child with no visual cues at all. When the time is right the language will come out and just keep flowing out.

5) I would be cautious about using any sign language. The problem is kids will use this as a crutch. The philosophy in oral training is to force our children to use their voice. To let them know that their voice has meaning. Let them know when they do use their voice that you understand, and always reward them with a huge amount of praise for using their voice.

The process is to give oral speech a chance before any signing is introduced. Some people will call signing a “bridge” to help facilitate oral speech. This may be true with some children but in general, I do not agree with signing. To take a child who has been implanted at or before 12 months of age and put them into a TC (Total Communication) or signing program in my opinion is medically wrong unless that child has other severe medical issues.

You may have professionals steer you into signing in the very beginning when you do not know better. BE CAUTIOUS. I would only encourage this if you want your child to know sign language with the possibility of their audition and articulation suffering. I would rather be the master of one modality than mediocre at two languages. The key here is that perfect spoken language and excellent audition is what we are concentrating on.

6) Keep names simple, one name for each item. Many items have numerous synonyms, but choose one name for each item. Stick with it until they fully understand that name. (A pair of shoes is SHOES, not sandals, boots, slippers, tennis shoes, etc. They are all shoes for the first six months. Same with plates, cups, etc.)

Make a list of 40 - 50 items and write down one name for each item and have the entire family use this one name. As time progresses and your child understands more, you can move onto the next synonym. Speak only one language to your child for the first three years post-implant. Do not confuse these children with a second language. In our case and because we do live in the United States, English is our primary language, Spanish our secondary language. These children can be excellent bilingual speakers but the exact mix needs to be researched. I did not want my daughter to be the test case of when to start a second language. There needs to be more research done on this particular subject; please check with your therapist.

We introduced Spanish at three years post-implant and went through the same hierarchy of language that was learned with English. One word and one name at a time. Overhearing conversation and having the opportunity to give input in a second language is also good. In our case I stick strongly to the three-year rule and would encourage others to do the same.

Full contextual language is very important also, and the key is to build a strong vocabulary and receptive language abilities with the parent and therapist explaining things in full context.

7) Play proper language-rich video tapes and DVDs. Let your child play with educational auditory toys that require good listening skills. These toys will ask a question and then your deaf child will need to make an appropriate response. Read every night if possible. Play children’s music CDs / tapes in the car, sing songs, and talk, talk, talk.
8) Turn on the implant or hearing aids within 30 minutes after your child wakes up. Keep them on all day long. Check the implant / hearing aids daily to make sure they are working properly. Know how these devices function and know how to troubleshoot and fix any problem that may arise. In general when it comes to cochlear implants they have a 10-year warranty on the surgically implanted device and three years on the external components. Listen to the microphone, make sure it is on the correct program, make sure the batteries are fully charged, and check for proper function every day. Check all wires for crimps, shorts, etc. Once a week use a Dry and Store dehumidifier unless you live in a humid climate, which may require use more often. Check with the manufacturer for proper care and maintenance.

9) Make sure your child has an appropriate program to learn oral speech and that the teaching environment has qualified professionals. Make sure the acoustics will accommodate learning with a noise-free classroom.

10) On weekends get out and explore the world. Go for a hike, a walk, go to the zoo, the grocery store, even around your neighborhood. Look at every item, explain what it is and give it a name. Make a fun game out of it even if you have other children—explain that we are building the vocabulary of our young deaf / HOH child. Not too much too fast, but just the right pace so your child can absorb every word and phrase.

The first five years of a child’s life are so crucial for proper language development. These things cannot wait and you just can’t start too early. As I did with my life and my daughter, this is a one-shot deal. I can set four to five years of my life aside and concentrate on my daughter because I have lived my life, I am an adult. What’s four or five years to me? Nothing. But for my daughter it is the most important time in her entire life. What is done now will shape her language forever. There will be a day in the next two to three years that you do not have to work as hard, you can relax because your child will be caught up with other children their age and you can treat them like a regular kid. It is not like this sacrifice has to be forever, just a few years. This author is willing to help with any advice or questions you may have.

Keep faith that your child will speak and listen and it will all pay off. Just ask me, it is a miracle all the things that our daughter can now do.

**Must Have Language Tools**

Here are a few must-have items that will help any child develop language.

1) Baby Einstein Video / DVDs. (Good language learning videos that children can watch.)

2) Videotapes of your child’s oral training therapies. Kids love to see themselves on TV. They can watch and learn over and over again.

3) LeapFrog Oral Language Toys and DVDs. (Buy toys that your child can learn from, not just take up time. These toys ask questions and need a response. Explain everything when playing together.) The DVDs will explain letters, phonetics and even simple math when your child gets a little bit older.

4) Picture books with clear colorful pictures. (So you can point to each picture and name it.)

5) The Ausplan book, this is a must-have book provided by Children’s Hospital Oakland’s Audiology department (510-428-3885). This book goes through every step of language development for cochlear-implanted children. You will refer to this book over the next three to five years and know exactly where your child is and what the next step will be in their habilitation process. You will be in sync with the
speech therapists and most importantly you will be able to know where your child is on the speech, language, and audition hierarchies.

These are the steps that a parent needs to take to give their child the best opportunity for success. You are in the driver’s seat, and at times, you may be in a position to need to disagree with the professional recommendations for treatment. You will feel confident that your viewpoint has merit because it has been researched and you have data to back it up. Keep in mind that an excellent working relationship with all parties concerned is so very important.

Do not be the one who sits back and is removed from this process. We all have jobs to do, families to care for, and homes to take care of. Ideally both mother and father should have similar knowledge in this process so they can talk about and bounce ideas and questions off each other.

In addition to the preceding list, you will need an experienced cochlear implant surgeon who has operated on hundreds of other children. If the experience is not there then demand a new surgeon. Do not be afraid to ask your doctor, “How many of these cases have you done? How many cochlear implant surgeries have you performed? What is the end result that you have experienced from other patients in this particular situation?” Definitely do not be rude or sound as if you know any more than they do. You need always to be gracious and polite. You will be dealing with the doctors and the other professionals for the next three to five years so do not get going on the wrong foot. Just be clear with what you want for your child. “I want my child to hear and speak very well. I have spoken to parents who have children that speak and hear great, and I want the same for my child. I will do whatever it takes to make this happen. We are committed as a family.” When your doctor or caregiver hears this they know you are on board. And believe it or not, you are one of the very special parents that really, really cares about the process and is committed to work with these fine people. They will bend over backwards for you as long as they know you are on board and you always show your sincere appreciation for what they do.

You also need an experienced pediatric audiologist who has diagnosed and mapped numerous children with their implants. Again do not be afraid to demand a different person if the experience is not present. Ask other people for referrals. In a lot of circumstances the audiologist is set up by the surgery center or sometimes the surgeon is recommended by the audiologist. Talk to other parents and find out what they think about a particular doctor or audiologist. Keep in mind you need a pediatric audiologist who has a great deal of experience with cochlear-implanted children, not a general practitioner.

Last but not least is how to pay for all of this fabulous care that your young son or daughter will be receiving. (This subject is covered more extensively in Chapter 7 of my book.) Here are my thoughts and experience in this area:

Most funding that you will need for the education of your child will be obtained from the text within the IDEA law. Please purchase a good book which covers the IDEA / IFSP / IEP process such as the Nolo Press series. The goals for your child and the services provided will be agreed upon at each IFSP /IEP meeting. Know what you want before each meeting, have professionals with you and be ready to back up every point as needed. Make oral education cost-effective for the folks holding the funding dollars. Our children can in many circumstances be mainstreamed by kindergarten or first grade with very little ongoing support. This is different from an ASL-taught child, who may need interpretive services throughout their school years and even into college.

Use the legal resource team at the Let Them Hear Foundation [www.letthemhear.org](http://www.letthemhear.org) to help fight any insurance denial for an implant. If you are forced to pay cash for any procedure, demand the same rate that the insurance company would pay the doctor. In many cases this could be 40% to 50% less than what they will charge a cash-paying customer. Ease the doctor’s mind by having cashier’s check(s) made out to each party and give these checks to the appropriate parties on or before the day of the uncovered procedure.
Put your entire life into this process for the next three to five years, look seriously at these guidelines, and know that educating yourself is one of the most important things in this entire process.

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