

Therapeutic Potential of Neurotrophin Gene Therapy in Noise-Induced Hearing Loss and Age-Related Neural Hearing Disorders

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Abstract

Neurotrophin (NT) cochlear gene therapy might perhaps give a single treatment that might greatly enhance neuronal survival, resulting in CI patients, provided the many challenges described above can be adequately addressed and safety concerns allayed by more animal model investigations. This is particularly crucial for juvenile CI patients, who have to rely on electrical hearing for the remainder of their lives, and whose outcomes are quite different. In addition, NT gene therapy may have the potential to treat patients with noise-induced hearing loss or neural presbycusis (e.g., age-related cochlear synaptopathy), where primary neuronal loss is a key cause of hearing loss. Animal research into noise-induced hearing loss has shown that even exposures that generate only reversible threshold alterations and no hair cell loss can lead to permanent loss of SGN synapses on hair cells, resulting in functional impairments and ultimately SGN degeneration. Cochlear synapses frequently precede both hair cell loss and threshold increases in human ears, according to current studies. Cochlear synaptopathy is characterized by ears with intact hair cell populations and normal audiograms as "hidden" hearing loss. Many frequent perceptual abnormalities, including speech-in-noise difficulties, tinnitus, and hyperacusis, are likely produced by suppressing affected neurons, which radically alters information processing. Thus, in the future, NT gene therapy may be successful in inducing SGN peripheral axon resprouting and synaptic regeneration into residual (or even regenerated) hair cell populations. We have demonstrated compelling evidence that, in this investigation, BDNF gene therapy can boost SGN survival and enhance peripheral axon maintenance or rerouting. NT-3 has been found in adult animals exposed to acoustic damage to induce synaptic regeneration of these fibers, reconnecting them to hair cells and their ribbon synapses, and restoring hearing function. Combining BDNF and NT-3 gene therapy may be the most effective way to maintain/restore a more normal cochlear neuronal substrate.

1. Introduction

Hearing loss frequently has negative educational, social, and vocational repercussions that have a substantial impact on one's quality of life. Furthermore, hearing loss is linked to cognitive impairment (Chern and Golub, 2019) and depression (Jayakody et al., 2018), particularly in the elderly. Although gene therapy options detailed in earlier papers in this Special Issue may improve these numbers in the future, for severe to profound hearing loss, the cochlear implant (CI) is now the standard of care. In general, today's CIs are quite successful. The average CI recipient utilizing the most up-to-date technology gets roughly 80% of high-context sentences right through voice recognition and can use a phone (Zeng et al., 2008). Music may even be enjoyed by the affluent (Drennan and Rubinstein, 2008; Won et al., 2010; Jiam et al., 2019). However, there is still a lot of variation in results among individual CI receivers (Firszt et al., 2004; Holden et al., 2013), especially in young people (Svirsky et al., 2000; Ortmann et al., 2017; Zhao et al., 2019), and many CI users get little or no benefit. The amount of surviving cochlear SGNs in individual CI recipients has been demonstrated to be a major factor determining performance, and this variability is likely connected to variances in auditory nerve survival (Kamakura and Nadol, 2016; Seyyedi et al., 2014). Furthermore, there is research that has found links between indirect functional markers of neural survival and CI results (Holden et al., 2013; Scheperle, 2017; Schwarts-Leyzac and Pfingst, 2018), bolstering the case for auditory nerve survival. As a result, there has been a lot of interest in recent years in looking into possible therapeutics for improving auditory nerve survival by slowing SGN degeneration following deafness. Osmotic pumps and other techniques have been employed in deafened animal models to administer a number of NTs directly to the cochlea in preclinical research, and the results have been encouraging, particularly with BDNF. However, due to concerns regarding infection and effectiveness duration, osmotic pumps are not a good alternative for extensive clinical use, and other techniques are not yet completely established (see Ma et al., 2019 for review). Recent research has focused on the idea of employing neurotrophin gene therapy to drive NT expression by cochlear target cells. This might pave the way for a one-time therapy that promotes long-term NT expression and better SGN survival following deafness.

This review first describes the data supporting exogenous NTs' effectiveness in supporting enhanced SGN survival within the cochlea following hearing loss in this review. We'll also go over the animal research that's been done so far with NT gene therapy in the inner ear and other non-auditory uses. Finally, we'll go through some of the remaining concerns, such as optimum vector selection, treatment schedule, and delivery location/method, among other things. Prior to contemplating clinical application, this must be settled.

Exogenous neurotrophins increase SGN survival following deafness. Over the last two decades, there has been a lot of interest in neurotrophic drugs that might promote SGN and auditory nerve survival and so improve CI results (see Staecker et al., 2010; Leake et al., 2013). The NTs, which are members of the nerve growth factor (NGF) family of proteins and comprise NGF, BDNF, neurotrophin-3 (NT-3) and NT-4/5, each of which binds to distinct high-affinity Trk family receptors, are of special importance. BDNF and NT-3, in particular, are known to play critical roles in SGN formation and maintenance. Neurotrophins control neuronal differentiation and survival throughout cochlear development (Farinas et al., 2001; Fritzsche et al., 1999; Rubel and Fritzsche, 2002; Yang et al., 2011). Hair cells, supporting cells of the organ of Corti, and neurons of the cochlear nucleus all offer neurotrophic support to SGNs (Fritzsche et al., 1999; Schecterson and Bothwell, 1994; Stankovic et al., 2004), and SGNs express the BDNF (TrkB) and NT-3 (TrkC) receptors (Schecterson and Bothwell, 1994; Ylikoski et al., 1993). Furthermore, BDNF and TrkB have recently been discovered in the developing human cochlea, indicating that they have a comparable role in SGNs in humans (Johnson Chacko et al., 2017). Both BDNF and NT-3 are involved in the maintenance of SGNs in the adult cochlea (Qun et al., 1999; Ylikoski et al., 1993), and the loss of this neurotrophic support after deafness leads to SGN degeneration via apoptotic cell death (Alam et al., 2007; Fritzsche et al., 1999). Furthermore, multiple investigations have shown that exogenous NTs administered directly to the cochlea through an osmotic pump over a period of weeks can preserve SGNs and enhance neuronal survival following deafness caused by different stressors (Ramekers et al., 2012; Leake et al., 2013; Ma et al., 2019). Deafened guinea pigs (Agterberg et al., 2008; Glueckert et al., 2008; Miller et al., 2007; Shepherd et al., 2008; Wise et al., 2005; Ramekers et al., 2015) and neonatally deafened cats (Agterberg et al., 2008; Glueckert et al., 2008; Miller (Leake et al., 2011). Other NTs, such as glial-cell-line-derived neurotrophic factor (GDNF) (Kanzaki et al., 2002; Maruyama et al., 2008; Yagi et al., 2000; Ylikoski et al., 1998) and Fibroblast growth factor (FGF), have been shown to have neurotrophic effects (Glueckert et al., 2008). Furthermore, while one study (Gillespie et al., 2003) reported rapid SGN loss after NT delivery was stopped, several other more recent studies (Agterberg et al., 2009; Leake et al., 2011; Shepherd et al., 2008) have shown that neurotrophic effects can last long after exogenous NT delivery is stopped (Agterberg et al., 2009; Leake et al., 2011; Shepherd et al., 2005; Leake et al., 2013). Importantly, when BDNF infusion was paired with CI implantation, a highly significant increase in SGN survival (>50 percent increase re: contralateral) was sustained when electrical stimulation from the CI was sustained 3–4 months after BDNF supply was stopped (Leake et al., 2013). Exogenous NT infusion has been shown in several labs to enhance the survival of radial nerve fibers in the osseous spiral lamina (the peripheral dendrites of the SGNs) as compared to deafened controls (Glueckert et al., 2008; Leake et al., 2011, 2013; Pettingill et al., 2007; Wise et al., 2005). Reduced thresholds and higher dynamic ranges for electrical stimulation provided by CI were related to improved fiber survival (Leake et al., 2013; Landry et al., 2013), which might enhance CI performance. BDNF injection, on the other hand, frequently results in widespread ectopic and chaotic sprouting of radial nerve fibers down into the scala tympani and spiraling hundreds of micrometers in the connective tissue encasing the implanted CI (Glueckert et al., 2008; Leake et al., 2011, 2013; Staecker et al., 1996). Ectopic fibers come in

both myelinated and unmyelinated forms, and they clearly show ectopic sprouting (i.e., normally, SGN peripheral axons never appear in the scala tympani, but are limited to the osseous spiral lamina and organ of Corti). Furthermore, even though these animals were given BDNF at one month of life, the structure of the cat's cochlea is mature at this age, and the sprouting fibers were still there when the mice were evaluated as young adults at around six months of age. Importantly, Glueckert et al. (2008) used immunolabeling after a combination of BDNF and FGF treatment to show that both the fibers within the osseous spiral lamina and the ectopic fibers elicited by BDNF treatment were afferent peripheral processes of SGNs, making them relevant to CI stimulation. However, efferent fiber survival was unaffected. Finally, electrophysiological recordings from the inferior colliculus in deafened, BDNF-treated animals have revealed that such sprouting can compromise the CI's optimal function by degrading the normally precise cochleotopic organization of the radial nerve fibers, and thus the selectivity of CI stimulation channels in the auditory midbrain (Leake et al., 2013). Recent developments in CI technology, including current focusing and virtual channel stimulation, rely on very spatially confined SGN activation, which would be harmed by sprouting.

3. Gene therapy for cochlear neurotrophin
Recent breakthroughs in cochlear molecular therapeutics, which have been addressed in numerous earlier papers in this Special Issue, are showing promising results in the development of therapeutic therapeutics for hearing loss by focusing on the correction of genetic abnormalities that cause hair cell loss. Preclinical studies have shown a potentially crucial role for NT treatment in reducing SGN degradation after deafness and enhancing Cis outcomes, as we've already said. Virally-mediated gene therapy has the benefit of needing only a single injection to induce safe and long-term expression of NTs by cells within the target tissue.

3.1. Rodent research

Several investigations in deafened mouse models have found that virally-mediated NT cochlear gene therapy employing a variety of NTs (BDNF, GDNF, NT3, CNTF, and others) reduces SGN degeneration and improves survival following damage. Ad vectors were used in the majority of previous research. Following cochlear administration of vectors pushing expression of NTs, notably BDNF, several investigations in deafened guinea pigs have observed higher SGN survival relative to contralateral after cochlear administration of vectors pushing expression of NTs, notably BDNF (Atkinson et al., 2012, 2014; Chikar et al., 2008; Nakaizumi et al., 2004; Rejali et al., 2007; Shibata et al., 2010; Wise et al., 2010, 2011).

Furthermore, research in additional deaf animal models, such as deaf mutant mice (Fukui et al., 2012) and rats exposed to blasts (Wu et al., 2011), has shown better SGN survival following virally-mediated NT delivery to the cochlea, confirming the cross-species hypothesis. After NT gene therapy, several research groups have observed better survival or signs of resprouting of the radial nerve fibers (the peripheral processes of the SGNs) (Atkinson et al., 2012, 2014; Chen et al., 2018; Fukui et al., 2012; Shibata et al., 2010; Wise et al., 2010). Because the fibers are closer to the CI electrodes, this might help CI function by lowering electrical stimulation thresholds and boosting spatial selectivity and temporal coding. Following cochlear NT gene therapy, certain studies have found indications of functional benefits with an implanted CI (Chikar et al., 2008; Budenz et al., 2015; Pfungst et al., 2017).

Most recent studies have used AAV vectors to drive expression of NT-3 and/or BDNF and have demonstrated excellent success in enhancing both SGN and radial fiber survival (Budenz et al., 2015; Pfingst et al., 2017; Chen et al., 2018). This switch to AAV was most likely motivated by the fact that AAV has been demonstrated to efficiently transport toxins to the inner ear while also being non-ototoxic (Ballana et al., 2008; Konishi et al., 2008; Lustig and Akil, 2012; Gyorgy et al., 2017; Pfingst et al., 2017; Suzuki et al., 2017; Tao et al., 2018). Furthermore, AAV has already been used in clinical trials with no negative consequences (see section 3, below). In deafened, implanted guinea pigs, Pfingst et al. (2017) documented the long-term (although varied) efficiency of AAV-mediated NT-3 gene therapy, as well as showing that psychophysical and electrophysiological techniques may be relevant for monitoring SGN density in the implanted cochlea. Budenz et al. (2015) found that while BDNF was more effective than NT-3 in preventing SGN degradation and improving long-term neural survival following deafness, NT-3 was more successful in triggering radial nerve fiber regrowth. These researchers believe that over-expression of both BDNF and NT-3 in combination may be the most effective way to improve overall neuronal survival.

3.2. Research on cats

The encouraging findings of NT gene therapy in deafened rodents prompted a new study to explore the possibilities for using gene therapy in the considerably bigger feline cochlea, bringing the results closer to the human cochlea (Leake et al., 2019). The animals were deafened as newborns before hearing onset (systemic neomycin injections) to imitate congenital deafness, which was a new component of this investigation. To allow for comparison with earlier studies in which exogenous BDNF was delivered by osmotic pumps at this age (Leake et al., 2011, 2013), and with the rationale that long-term SGN survival and improved CI outcomes are especially important for the pediatric population, gene therapy was administered when the animals were about a month old. AAV2 encoding for BDNF (under control of the CGA promotor) and AAV5-GDNF, both of which have proven effectiveness in previous systems, were compared (CBA promotor). At 3 months after injection, both vectors had minor neurotrophic benefits, with around 6% of the normal SGN population saved compared to the contralateral.

GDNF expression, on the other hand, resulted in undesired fiber sprouting into the scala tympani and did not enhance the number of surviving fibers inside the osseous spiral lamina. In comparison to untreated ears, AAV2-mediated BDNF expression resulted in more than twice the number of surviving radial nerve fibers, with no ectopic or disordered sprouting. Following up on the positive results of AAV2-BDNF, researchers wanted to see if the neurotrophic benefits would last when the post-injection survival time was prolonged to 6 months. Significant neurotrophic benefits were shown in this long-term investigation, with increased SGN neuronal survival maintained throughout the cochlea compared to the contralateral reference. Six months following AAV2-BDNF injections, total mean SGN survival was 53% normal vs. 39% contralateral, indicating that around 14% of the normal SGN population was rescued. When expressed as a percentage increase adjusted to contralateral survival, this translates to a 35 percent increase in SGN survival compared to the control group. It's worth noting that SGN survival in these early-deafened cats was projected to be around 75% of normal at the time of viral injections (1 month of age) (Leake et al., 2011). Despite the fact that AAV2-BDNF evoked a strong neurotrophic effect, the SGNs in the injected ears continued to degenerate. Transfection happens quickly, but only a small number of cells are transduced, and maximal NT expression appears to take significantly longer, according to immunohistochemistry done two weeks after viral injections. The survival of radial nerve fibers in these deafened animals was also measured by counting them in sections cut orthogonal to the radial plane at three different cochlear sites. SGN peripheral fiber survival was consistently greater in the injected ears in all three cochlear regions compared to the contralateral in the 6-month AAV2-BDNF group. After AAV2-BDNF therapy, fiber survival averaged 47 percent of normal, which was twice the value recorded on the opposite side (24 percent of normal).

The findings in a variety of deafened animal models utilizing a variety of viral vectors and NTs delivered to the cochlea imply that NT cochlear gene therapy might be a feasible technique for reducing SGN and radial nerve fiber degeneration after deafness. As a result, a therapy that just requires a single injection has a lot of potential for fostering long-term development in the cochlear neural substrate and, as a result, improving CI results.

3.3 NT gene therapy for cochlear synaptopathy

Animal studies of noise-induced hearing loss have shown that exposures causing only reversible threshold shifts and no hair cell loss can result in permanent loss of SGN synapses on IHCs (cochlear synaptopathy), which can lead to functional deficits and eventually SGN degeneration when followed long-term (Kujawa and Liberman, 2009). Furthermore, multiple investigations have indicated that NT-3 can preserve or even repair IHC synapses, allowing hearing function to be restored after sonic stress (Sly et al., 2016; Suzuki et al., 2016; Wan et al., 2014; Wang et al., 2011). As a result, it's worth noting that AAV-mediated NT-3 overexpression can likewise protect against and heal noise-induced cochlear synaptopathy, according to recent animal research (Chen et al., 2018; Hashimoto et al., 2019).

Furthermore, new research in both noise-exposed and aging human ears has revealed that cochlear synapse degeneration frequently occurs before both hair cell loss and threshold increases (Sereyenko et al., 2013; Kujawa and Liberman, 2015; Liberman, 2015, 2017; Liberman and Kujawa, 2017). Many common perceptual anomalies, including speech-in-noise issues, tinnitus, and hyperacusis, are likely caused by the silencing of afflicted neurons, which drastically changes information processing. As a result, NT gene therapy may one day be effective for prompting SGN peripheral axon resprouting and synaptic regeneration to innervate residual (or regenerated) hair cell populations in situations of "hidden" hearing loss.

4. Non-auditory applications of neurotrophin gene therapy

NTs and members of the neurotrophin family serve crucial roles in the formation, maintenance, and repair of the central nervous system (CNS), much as they do in the auditory system (see Review by Huang and Reichardt, 2001). Several gene therapy approaches have been investigated in animal models and clinical trials of several neurodegenerative diseases, including retinal and optic nerve degeneration, spinal cord injury, Alzheimer's disease, Parkinson's disease, Huntington's disease, and amyotrophic lateral sclerosis, over the last three decades (see Review by Blesch et al., 1998; Khalin et al., 2015; Daly et al., 2018; Hardcastle et al., 2018; Hodgetts and Harvey, 2017; Mestre and Sampaio, 2017). Although significant progress has been made, and mounting evidence from animal studies supports the efficacy of neurotrophin gene therapy in preventing neuronal degeneration and promoting neural repair, a number of technical challenges must be overcome before these applications can be successfully applied to human patients. This section focuses on the efficacy of various gene therapy applications as well as the current problems they face. The following discussion outlines our current understanding of how to improve delivery protocols to improve the effectiveness and specificity of NT gene therapy in various neurodegenerative conditions, as well as the implications for possible use in auditory nerve degeneration and hearing loss.

4.1. Neurodegenerative diseases that aren't Alzheimer's

Gene therapy (such as intravitreal gene transfer) has been a popular treatment option for hereditary optic neuropathies and other retinal illnesses due to the retinal ganglion cells' anatomical accessibility (Thanos and Emerich, 2005; Yu-Wai-Man et al., 2014). Although the first results are promising, these gene therapy techniques are still in the early stages of research, and further proof of their efficacy, specificity, and safety is needed before they are applied to human patients. NGF, BDNF, and NT3 are members of the NGF family of NTs, which play important roles in retinal ganglion cell survival. For example, retinal ganglion cells produce BDNF locally (Herzog and von Bartheld, 1998). In various animal models of retinal degeneration, GDNF, a member of the transforming growth factor superfamily, protects the retina (Thanos and Emerich, 2005).

Previous research has suggested that NT deficiency and dysfunction have a role in the etiology of glaucoma (Bringmann et al., 2006; Johnson et al., 2011). In a rat model of optic nerve transection, AV vectors were used to increase Müller glial expression of BDNF (Di Polo et al., 1998). AAV-mediated BDNF transfer to Müller cells protected photoreceptors from light-induced retinal degeneration, according to Gauthier et al. (2005). AAV-GDNF reduced photoreceptor loss for at least 45 days in an animal model of human retinitis pigmentosa (S334ter-4 rhodopsin transgenic rat) using AAV-mediated gene delivery (McGee Sanftner et al., 2001). These findings show that increasing endogenous retinal synthesis of specific NTs via viral vectors might reduce or prevent retinal and optic nerve degeneration.

BDNF is the most studied NT protein in experimental spinal cord injury (SCI), and it is involved in axonal sprouting, neuroprotection, myelination, adaptive synaptic plasticity, synaptic transmission, and antioxidative actions (Kovalchuk et al., 2004; Weishaupt et al., 2012). In a rat model of total spinal cord transection treated with local BDNF overexpression induced by AAV1/2 vectors under control of the neuron-specific human SYN 1 promoter, Ziemińska et al. (2014) showed a considerable increase in treadmill locomotor capacities. Within 30 minutes following spinal cord transection, AAV-BDNF intraspinal injections were given, and locomotor function improvement was shown as early as two weeks following therapy and sustained for at least seven weeks. In the AAV-BDNF-treated rats, there was an increase in excitatory neurotransmission-related molecules. Around the damaged region, however, an altered balance of excitatory and inhibitory neuronal activity was found. These modifications might have had a role in the motoneuron hyperexcitability that was common following injury. A retrograde adenovirus (Ad-mediated) BDNF gene transfection resulted in decreased cell death in neurons and oligodendrocytes in a chronic SCI model (twy/twy mice) (Uchida et al., 2012). This neuroprotective effect was observed four weeks after Ad-BDNF administration, but the treated mice were not functionally evaluated. In rat models of spinal cord injury, two previous investigations found that virus-mediated BDNF gene transfection decreased neural cell death and improved axonal regeneration and locomotor functional recovery for at least 6 weeks after injections (Koda et al., 2004; Nakajima et al., 2010). Although AAV-mediated BDNF + GDNF transfection improved motor neuron survival, it did not result in functional recovery following ventral root avulsion (Blits et al., 2003). Martínez-Gálvez et al. (2016) used a cervical spinal cord injury model to show that delivering AAV7-mediated gene transduction for TrkB, a high-affinity receptor for BDNF, improved respiratory function. Research has looked at the efficacy of NT gene therapy employing various members of the NT family, combinations of various NTs, or coupled NT overexpression with cell-based treatment (e.g., stem cell or Schwann cell implants) (see review by Blesch et al., 1998; Hendriks et al., 2004; Harvey et al., 2015; Hodgetts and Harvey, 2017). The injection of an AAV-BDNF and AAV-NT-3 combination into the gray matter of the spinal cord, caudal to a Schwann cell graft implanted in damaged rats, improved locomotor performance, but no evidence of axon regrowth from the Schwann cell implant was seen (Blits et al., 2003). It's worth noting that in all of the experiments described above, virus-mediated gene therapy was delivered immediately or shortly after damage (e.g., within 3 days).

In neuropathological illnesses such as Alzheimer's disease, Parkinson's disease, Huntington's disease, and schizophrenia, dysregulation of NTs such as BDNF and NGF is a crucial component (Phillips et al., 1990; Sampaio et al., 2017; Simmons, 2017). Alzheimer's disease is a neurological illness that progresses over time and is the leading cause of dementia in older people (see review by Loera-Valencia et al., 2019). Reduced NT expression and deregulation of their relevant signaling pathways have been linked to Alzheimer's disease and Parkinson's disease, according to studies (Alves et al., 2016; Ventriglia et al., 2013). Several animal studies (Nagahara et al., 2009, 2013; Alves et al., 2016; Jiao et al., 2016) as well as clinical trials have revealed varying levels of effectiveness in NT gene therapy employing a viral-based delivery strategy (Rafii et al., 2014; Tuszynski et al., 2015; Malkki, 2015). The injection of lentivirus-mediated BDNF into the cortex of APP amyloid-transgenic mice, an animal model of Alzheimer's disease, increased synaptic protein production, reversed synapse degradation, and restored learning and memory performance (Nagahara et al., 2009). A comparable lentivirus-mediated BDNF therapy was given to both elderly rats and primates in this investigation. 2–4 weeks following therapy, these rats showed improved cognitive performance as well as decreased neuronal atrophy. However, the positive neuroprotective impact was not reliant on the presence of amyloid plaques. Jiao et al. (2016) used BDNF gene therapy in P301L mice, an Alzheimer's disease animal model characterized by age-related tau pathology and memory loss. Before severe tau pathology and cognitive impairment, the mice were given intraventricular injections of AAV8-BDNF at 3 months of age. 9 months following therapy, neuronal structures were restored and cognitive function improved, but there were no improvements in the tau pathogenic status.

Another key NT is NGF, which is required to prevent the loss of basal forebrain cholinergic neurons, which are commonly lost early in Alzheimer's disease. Several Phase I clinical trials have revealed that virus-mediated NGF treatment might prevent or minimize cholinergic neuron degeneration in Alzheimer's disease patients. In one study, AAV2-NGF was injected bilaterally into the basal forebrain areas of ten Alzheimer's disease patients (Rafii et al., 2014). All of the patients' brains showed a trophic response to NGF transduction, as well as a slower pace of cognitive deterioration. The bioactivity of AAV2-mediated NGF expression was discovered in brain autopsy tissues in this work; the longest post-treatment evaluation time was roughly 24 months, indicating that this technique may be practical and capable of creating reasonably long-term and physiologically active NT expression. NGF was previously given by autologous fibroblasts transfected with NGF-leukemia viruses and transplanted into the basal forebrain area containing cholinergic neurons (Tuszynski et al., 2005). Over the course of a two-year monitoring period, this *ex vivo* investigation found structural and functional improvements. The same group's second phase I investigation found that deteriorated neurons responded to NGF gene therapy in all patients (Tuszynski et al., 2015). The participants in this research were followed for periods ranging from 11 months to ten years. These findings imply that *ex vivo* or *in vivo* administration of the virus-mediated neurotrophin gene might be used to treat Alzheimer's disease and other neurodegenerative illnesses.

4.2. In non-auditory systems, lessons learnt from NT gene therapy research

A rising body of research suggests a relationship between age-related deafness, blindness, and dementia (see reviews by Mancino et al., 2018; Chern and Golub, 2019), pointing to shared processes of neurodegeneration in the peripheral auditory nerve, retinal and optic nerve, and the brain. It's also plausible that the loss of sensory information and social contact has a negative impact on neurocognitive function, hastening cognitive decline. Recent research into cochlear implantation in older people has found improvements in attention and working memory, such as with the operation span task, as evidence for this concept (Volter et al., 2018). A better understanding of the challenges and issues identified in studies of NT gene therapy in retinal and optical nerve degeneration, spinal cord injury, Alzheimer's disease, and other neurodegenerative disorders has significant implications for the design and refinement of investigations into these applications in the auditory system and hearing loss. Three important challenges are explored here in order to better understand how to improve the efficacy of NT gene therapy techniques. First, the efficacy of NT gene therapy is very time-dependent, with the better outcome occurring earlier in the illness stage when treatment is initiated. The therapies were administered either immediately or shortly after damage in most of these animal experiments, as described above under CNS trauma such as spinal cord damage. Similarly, late-stage patients with Alzheimer's disease and other neurodegenerative disorders had little response to neurotrophic gene therapy (Bartus and Johnson, 2017b). A better understanding of the dynamic changes in endogenous NF expression features in different populations of neural cells and conditions of inflammation after CNS injury or at different stages of neurodegenerative disease would be beneficial. Second, determining the appropriate places to inject gene therapy reagents is crucial. Because the method may deliver regulated, long-term physiologically active NTs in the targeted areas or cells, virus-mediated gene therapy has emerged as a viable therapeutic strategy for SCI, Alzheimer's disease, and other neurodegenerative diseases. For many of these uses, however, creating an effective and safe administration method remains a difficulty. For example, the *ex vivo* delivery strategy for NGF gene transduction in the cholinergic neurons of the basal forebrain area was developed in the phase I clinical trial of Alzheimer's disease stated above (Tuszynski et al., 2005). Finally, more research is needed to discover the optimum method for administering the medication and if a certain quantity or amount of virus is sufficient (Harvey et al., 2015; Bartus and Johnson, 2017a, 2017b). Potential consequences of exogenous neurotrophin gene expression on the function of host neural cells, such as neurotransmitter dysregulation or impairment of brain plasticity, must be carefully considered. For example, in some situations of spinal cord damage, overdosing on neurotrophic factors may result in adverse effects such as increased sensitivity to pain or seizures (Cunha et al., 2009; Weishaupt et al., 2012; Hodgetts and Harvey, 2017). Negative feedback of the BDNF/TrkB signaling pathway and 'trapping' of regenerated axons results from continued high levels of BDNF expression (Eaton et al., 2002; Blits, 2003).

5. Clinical application future aims and obstacles

Preclinical cochlear gene therapy research has focused on the preservation of SGNs in order to either preserve/restore hearing or improve the performance of cochlear implants. As previously mentioned, multiple studies have shown that Ad or AAV-mediated delivery of NTs (BDNF, GDNF, and NT-3) can help to prevent cochlear SGN degeneration following deafness. Cochlear gene therapy, unlike traditional pharmacological therapies, is a complicated biological treatment whose success is dependent on a variety of circumstances. These include, but are not limited to: 1) viral vector design optimization (safety, toxicity, and target specificity) and high quality vector production (efficacy at the lowest possible dose; ability to replicate long-lasting and stable levels of expression); 2) viral delivery site and method; and 3) intervention at an appropriate stage of the hearing disorder.

5.1. Choosing the best viral vector (s)

Previous gene therapy research has documented a number of modified replication-deficient viral vectors, including the Ad and AAV vectors, which have emerged as the most extensively employed tools for virally-mediated transport of NTs to the cochlea.

Adenovirus (Adenovirus) (Adenovirus) (Adenovirus)

In the last two decades, ad has become a popular candidate for gene therapy (Lee et al., 2017). As a result, it's no surprise that it's one of the two main viral vectors utilized in cochlear gene therapy (Praetorius et al., 2009; Staecker et al., 2014). Ads have a relatively large cloning capacity of more than 10 kb and have tropism for a variety of cochlear cell types, which are two benefits for gene delivery. Because many illnesses are caused by mutations in genes with coding sequences that exceed the AAV capacity, the cloning capacity is critical. As a result, ad vectors have a significant advantage over AAV in terms of cloning capacity. The length of transgenic expression is generally a few weeks to months, which is beneficial for applications that need a limited time for expression. This is also one of its major drawbacks for NT gene therapy or hereditary causes of hearing loss, where long-term gene expression is critical. Another drawback of the ad is its proclivity for eliciting an immunological response. Although newer generations of ads have been developed to lessen this risk, if repeated administration is necessary, immune-related adverse effects may be exacerbated. Despite their great cloning capability, the more intricate structure of Ad and the higher potential for eliciting undesired immune responses may restrict their use in the ear.

5.1.2. Adeno-associated virus

AAV as a gene therapy vector has been used in several studies to partially or completely restore hearing loss in mouse models of hereditary deafness (Akil et al., 2012; Askew et al., 2015; Emptoz et al., 2017; Geng et al., 2017; Isgrig et al., 2017; Pan et al., 2017; Akil et al., 2019a, b; see other papers in this Special Issue). AAV vectors have the capacity to effectively transduce cells that have completed mitosis (Colella et al., 2018). They also offer superior safety profiles (low immunogenicity), which gives them a leg up over ads. Furthermore, it has been established that AAV is not ototoxic (Ballana et al., 2008; Konishi et al., 2008; Lustig and Akil, 2012; Gyorgy et al., 2017; Pflingst et al., 2017; Suzuki et al., 2017; Tao et al., 2018). Importantly, AAV vectors have been shown to be effective for gene delivery to the eye and other organs in proof-of-concept investigations and clinical trials (Simonelli et al., 2010; Flotte et al., 2011; Nathwani et al., 2011; Bowles et al., 2012; Le Meur et al., 2018). Furthermore, non-replicating recombinant AAV vectors may efficiently deliver transgenes to a variety of cochlear cell types, including non-dividing neurons and hair cells. The virus does not integrate into the host genome; instead, it stays episomal, resulting in stable, long-term transgenic expression (Xia et al., 2012). Because long-term expression is vital for many human applications, AAV vectors' long-term expression provides a significant advantage over Ad vectors. Because NT expression is likely to be necessary in the long run, AAV vectors may be the best alternative for delivering NTs to the inner ear. Finally, the cochlea is a good candidate for gene transfer because it is separated from the rest of the body, reducing viral propagation and immune system exposure.

5.1.3. Serotypes and promoters

Tissue-specific promoters can be used to achieve specificity in gene delivery. Although using a cochlear cell-specific promoter to guide transgene expression may reduce undesirable off-target effects, it is uncertain if the promoter truly improves cochlear cell transduction efficiency. In other tissues, such as the heart (Ai et al., 2008; Merentie et al., 2016), toxicity and inflammation have been observed with widely active promoters (CMV, CBA, etc.) but not with cell-type-specific promoters (Klein et al., 2006; Watakabe et al., 2015). One possible explanation for this toxicity is because widely active promoters generate greater transgenic expression than cell-type-specific promoters. The use of cell-specific promoters can limit transgenic expression in non-target cells and improve gene delivery specificity to the cell type of interest. There are a variety of potential cochlear cell-type-specific promoter options to choose from. Myosin VIIA promoter, elongation factor 1 promoter, neuron-specific enolase promoter, and glial fibrillary acidic protein promoter are examples of such promoters. These promoters have been cloned and studied in detail. The myosin VIIA promoter, which is expressed strongly and selectively in the hair cells of the cochlea and vestibule, was identified by Boeda et al. (2001).

Following cochlear injection in adult rats and mice, Lui et al. (2007) found that the myosin VIIA promoter generated selective expression of eGFP inside hair cells. The neuron-specific enolase promoter and the elongation factor 1 promoter both induced selective eGFP expression inside SGN and spiral ligament cells, according to these same researchers (Lui et al., 2007). Rio et al. (2002) had previously discovered glial fibrillary acid protein promoter selectivity in all cochlear supporting cells shortly after birth.

Transfection specificity can also be achieved by retargeting AAV to distinct cellular receptors or utilizing various AAV serotypes with various binding sites (Nam et al., 2011). The vitality of the cochlear cells targeted for viral transfection is expected to determine the efficacy of NT gene therapy. When the organ of Corti (OC) is chosen as the target (Wise et al., 2010), continuing degradation due to hearing disease may restrict the ability of NT gene therapy to offer the neurotrophic support required to protect SGNs. Even after significant OC degradation, viral transfection of cells inside the scale medium was still achievable (Wise et al., 2011). Supporting cells (e.g., pillar and Deiters' cells), cells inside the stria vascularis and spiral ligament of the cochlear lateral wall, endosteal cells covering the scala compartments, and interdental cells inside the spiral limbus are all examples of these cells (Wise et al., 2011). The AAV serotype, viral load, and promoter combinations that successfully transduce various auditory cell types are largely understood. Therefore, optimizing transduction efficiency is critical to enhancing the therapeutic impact of NT gene therapy. Various AAV serotypes, including AAV2 (Budenz et al., 2015; Pflingst et al., 2017; Leake et al., 2019), AAV8 (Chen et al., 2018), and AAV5 (Chen et al., 2019), have been employed for neurotrophin administration in animal investigations (Leake et al., 2019). Despite the fact that all of these studies showed an increase in SGN survival after deafness, there is no direct comparison of the efficacy of the various AAV serotypes used for NT gene therapy due to differences in other variables such as animal models, delivery sites and methods, virus concentration and dose, and so on. Kilpatrick et al. (2011) studied the transduction effectiveness and cellular specificity of multiple AAV vectors (serotypes 1, 2, 5, 6, and 8) in normal and drug-deafened ears in a single research. This study found that all five AAV serotypes effectively transduced the above-mentioned common cochlear cell types, implying that any of the AAV serotypes may be utilized for effective NT SGN gene therapy. The recombinant AAV2 serotype, on the other hand, appears to be the best option for two reasons: 1) It effectively transduces the cochlear cells that are being targeted. 2) It is the most often utilized viral vector in clinical trials for other organs (e.g., ocular gene therapy), and it is now being utilized to treat Leber's congenital amaurosis (Cideciyan et al., 2013; Bainbridge et al., 2015; Russell et al., 2017) and choroideremia (Cideciyan et al., 2017). (MacLaren et al. 2014; Edwards et al. 2016).

To date, research has shown that for each viral design, sensitive tests must be developed that are particular to the organ and cell types being targeted. Such tests will allow the development of vectors that may be safely used to deliver optimal dosages of vectors, potentially improving both safety and effectiveness (Xiong et al., 2019). If safer vectors can be developed, a larger number of cochlear cells could be transduced, resulting in increased effectiveness and fewer safety issues.

5.2. Determining the optimal concentration/dosage; clinical consequences

While inner ear gene therapy may prove to be a beneficial therapeutic technique in the future, safety concerns about viral gene delivery must be carefully examined in relation to the procedure's predicted advantages. It will be important to discover a viral titer that would offer the intended benefit while provoking minimum or no harm in order to optimize virus-mediated gene therapy for clinical use. The viral dosage and the promoter are the two parameters that influence a viral vector's transduction effectiveness, and they also have the strongest link to toxicity. Other factors are likely to have a role in toxicity as well (e.g., stocks with a high degree of endotoxin, or non-viral protein contamination). This emphasizes the need for tailoring the viral load and promoter design in the development of cochlear-targeted gene therapy in order to maximize the therapeutic impact.

5.2.1. Dose of vector

Current cochlear local delivery methods can result in just a small fraction of target cells being transfected since they are close to the injection site. A more thorough infection would almost certainly increase the intended effect (S), but it would necessitate a higher viral load, which might cause toxicity. In animal investigations, toxicity associated with greater dosages has been shown in the eyes and other organs (Mingozzi and High, 2013; Hinderer et al., 2018; Vandenberghe et al., 2011; Ramachandran et al., 2017; Khabou et al., 2018). According to Akil et al., 2019a, the overexpression of hGDNF induced by AAV5-hGDNF in newborn mice resulted in severe neurological symptoms and hearing loss due to Purkinje cell loss and cochlear nucleus pathology. Extremely high levels of transgenic protein expression should thus be avoided, especially for proteins with potential neurological roles (Akil et al., 2019a). This can be accomplished by lowering the amount of virus injected into the ear to reduce toxicity while still ensuring that the protein of interest is expressed at a high level in the targeted cells.

5.3. Scala tympani vs. Scala media as a route to administration.

Various gene delivery pathways have been successfully applied to sensory hair cells, spiral ganglion neurons, and cells in the stria vascularis in a variety of animal experiments (see Lustig and Akil, 2012; Géléoc and Holt, 2009; Chien et al., 2015; Ma et al., 2019). The surgical delivery to the inner ear space, which must limit harm to the inner ear while maximizing viral transduction of the target cells across the cochlea while retaining hearing function, is perhaps the most predictable of all the factors. The dosage delivered to the target cells may be reduced if the vector is reabsorbed outside of the cochlea. In addition, the injected vector particles may trigger an immune response against the viral capsids, reducing the quantity of effective vector particles in the cochlear region.

For effective targeting of cochlear cells, a safe and repeatable delivery of gene therapy vector into the inner ear is required. The comparison here is between two major delivery routes: 1) into the perilymph of the scala tympani, and 2) into the endolymph of the scala media. The most common approaches to cochlear gene therapy have been scala tympani operations, which can be done through the round window membrane, oval window, or direct cochleostomy via the bone otic capsule. Among them, the route through the round window membrane has been recognized as the best strategy for conserving residual hearing, and multiple investigations have shown that NT gene therapy using this technique is successful. However, as compared to the scala media technique, the scala tympani technique demonstrated reduced transduction effectiveness in cochlear cells. A cochleostomy via the cochlear lateral wall or a direct injection through the basilar membrane are used to reach the scala media pathway (Shibata et al., 2009; Wise et al., 2010; Kilpatrick et al., 2011; Chang et al., 2015). Due to the intricacy of the organ of Corti and the significance of the endolymphatic barrier and ion homeostasis for optimal hearing function, surgical operations for this pathway are more likely to cause hair cell injury and hearing loss. Kilpatrick et al. (2011) described an effective AAV inoculation method in mice's ears using the scala medium with little injection stress. The quantity (350 nl) and pace of injection into the scala media via the cochlear lateral wall were accurately controlled in this work using a microinjection device (WPI) capable of delivering quantities in the nanoliter (nl) to microliter (l) range. In adult mice deafened with kanamycin and furosemide, this method resulted in high AAV8 transduction efficiency in the auditory nerve. Finally, the best injection site will be determined by the precise cells that the gene therapy is targeting. The scala tympani, for example, is likely to be a superior choice if the SGNs are the target cells since it is right near to Rosenthal's canal, whereas the scala medium is considerably further away from the ganglion. Furthermore, strong junctional complexes across all cells isolate the scala media from the rest of the cochlear structures, restricting this specific fluid region.

5.4. Administration timing; consideration of the host cochlear milieu, glial cell activation, and post-deafness inflammatory reactions

The time of the therapeutic agent's delivery is a critical component in neurotrophin gene therapy's success. Gene therapy research, including Alzheimer's disease and CNS damage such as spinal cord damage, as discussed above, implies that the earlier the treatment (e.g., gene therapy administered soon after damage), the better the results (Harvey et al., 2015; Bartus and Johnson, 2017a, 2017b). Early treatment is thought to reduce axonal retraction, preserve myelin, and modulate the phenotype and activation of microglia/macrophages recruited from the circulation. Loss of the auditory nerve occurs in the peripheral auditory nervous system as a consequence of either primary or secondary degeneration after hair cell loss as a consequence of cochlear insults (Spoendlin, 1984; Leake and Hradek, 1988; Kujawa and Liberman, 2009; Lang, 2015a, b; Liberman, 2015).

Following cochlear insult caused by noise exposure and the administration of ototoxic medications, many pathological changes emerge in non-neuronal cells of the damaged auditory nerve. Demyelination, disruption of the Ranvier node, and activation of glial cells, as well as enhanced macrophage recruitment and activation, are among the pathogenic alterations (or other immune cells). Importantly, as observed in multiple animal investigations, all of these pathogenic changes occur in a time-dependent sequence (Lang et al., 2011, 2015; Tagoe et al., 2014; Panganiban et al., 2018; Kohrman et al., 2019). Previous research has revealed that in order for the NT treatment to be successful, it must be used in the early stages of illness, when there are still enough healthy neurons to react to treatment (Harvey et al., 2015; Quintino et al., 2019). However, the onset of SGN degenerative changes differs substantially between animal models (Wise et al., 2011; Leake et al., 2019). In applications of NT gene therapy, careful timing of administration should be included in the experimental design, based on consideration of the cochlear pathophysiological circumstances following deafness, in particular, the pathologies of these non-neuronal cells.

5.5. Administration site: endogenous cochlear NT expression tonotopic gradients

Both physically and functionally, the neurotrophins BDNF and NT3 play critical roles in the formation and early maintenance of the auditory nerve's tonotopic organization (Pirvola et al., 1992; Davis, 2003; Flores-Otero et al., 2007; Fritzsche et al., 2015). The *Bdnf* gene knockout produces a considerable loss of basal turn spiral ganglion neurons, whereas the NT-3 gene knockout produces a considerable loss of auditory nerve innervation to hair cells in the apical turn (Fritzsche et al., 1997). Furthermore, in both the postnatal and adult auditory nerves, BDNF has a greater expression in the basal area, but NT-3 appears to have a larger concentration in the apex (Farias et al., 2001; Sugawara et al., 2007). Adamson et al. (2002) used a unique in vitro setup to show that BDNF and NT3 had opposing impacts on SGN firing patterns. In particular, BDNF increased the activity of the SGNs in the apex, but had no effect on the neurons in the basal turn. In contrast, when neurons in the apex and base were exposed to NT-3, their firing patterns resembled those of the apical control. It is critical to evaluate the right location and cells in the cochlea that the gene therapy would target, similar to the present issues we outlined above in the research of spinal cord damage and Alzheimer's disease. When developing innovative therapeutic uses of NT gene therapy, it may be critical to consider how to minimize or prevent disrupting the tonotopic firing features of the surviving SGNs. The gene therapy must be tailored to maintain/restore the normal intrinsic firing properties of the SGN in order to achieve the long-term objective of restoring hearing through hair cell regeneration or attaching surviving hair cells to the auditory nerve. On the other hand, inherent firing features of SGNs may be overpowered by the more crude direct electrical stimulation supplied by a CI, and hence may not be as significant in the case of prospective use of NT gene therapy to sustain increased survival of SGN for application of a CI.

6. Conclusions

NT cochlear gene therapy might possibly provide a single treatment that might dramatically improve neuronal survival and results in CI patients if the multiple problems listed above can be appropriately addressed and safety concerns allayed by additional study in animal models. This is especially critical for pediatric CI patients, who must rely on electrical hearing for the rest of their lives and whose results are very diverse (Svirsky et al., 2000; Ortmann et al., 2017; Zhao et al., 2019).

Furthermore, in the future, NT gene therapy may have the potential to help individuals with noise-induced hearing loss or neural presbycusis (e.g., age-related cochlear synaptopathy), in whom primary neuronal loss is a major cause of hearing loss. Animal investigations of noise-induced hearing loss have demonstrated that even exposures that cause only reversible threshold changes and no hair cell loss can result in permanent loss of SGN synapses on hair cells, resulting in functional deficiencies and eventually SGN degeneration (Kujawa and Liberman, 2009). Degeneration of cochlear synapses often precedes both hair cell loss and threshold increases in human ears, according to current studies (Sereyenko et al., 2013; Kujawa and Liberman, 2015; Liberman, 2015, 2017). Cochlear synaptopathy has been described as "hidden" hearing loss in ears with intact hair cell populations and normal audiograms (Schaeffe and McAlpine, 2011). Many common perceptual anomalies, including speech-in-noise issues, tinnitus, and hyperacusis, are likely caused by the silencing of afflicted neurons, which changes information processing dramatically. Thus, NT gene therapy may be effective in the future in prompting SGN peripheral axon resprouting and synaptic regeneration to innervate residual (or even regenerated) hair cell populations. We have shown strong evidence that BDNF gene therapy can increase SGN survival and stimulate maintenance or resprouting of peripheral axons in this study. In adult animals exposed to acoustic damage, NT-3 has been found to induce synaptic regeneration of these fibers, reconnecting them to the hair cells and their ribbon synapses, as well as to restore hearing function (Sly et al., 2016; Suzuki et al., 2016; Wan et al., 2014; Wang et al., 2011; Hashimimoto et al., 2019). As previously proposed by Budenz et al. (2015), combining BDNF and NT-3 gene therapy may be the most effective way to maintain/restore a more normal cochlear neuronal substrate.

References

- Adamson CL, Reid MA, Davis RL, 2002. Opposite actions of brain-derived neurotrophic factor and neurotrophin-3 on firing features and ion channel composition of murine spiral ganglion neurons. *J. Neurosci* 22 (4), 1385–1396.
- Agterberg MJ, Versnel H, de Groot JC, Smoorenburg GF, Albers FW, Klis SF, 2008. Morphological changes in spiral ganglion cells after intracochlear application of brain-derived neurotrophic factor in deafened Guinea pigs. *Hear. Res* 244, 25–34. 10.1016/j.heares.2008.07.004.
- Agterberg MJH, Versnel H, van Dijk LM, de Groot JC, Klis SF, 2009. Enhanced survival of spiral ganglion cells after cessation of treatment with brain-derived neurotrophic factor in deafened Guinea pigs. *J Assoc Res Otolaryngol* 10, 355–367. 10.1007/s10162-009-0170-2.
- Abdelhamid, H. N., M. Dowaidar, M. Hällbrink, and Ü. Langel. 2019. Cell Penetrating Peptides-Hierarchical Porous Zeolitic Imidazolate Frameworks Nanoparticles: An Efficient Gene Delivery Platform. *SSRN Electron. J.* https://papers.ssrn.com/sol3/papers.cfm?abstract_id=3435895.
- Abdelhamid, Hani Nasser, Moataz Dowaidar, and Ülo Langel. 2020. Carbonized Chitosan Encapsulated Hierarchical Porous Zeolitic Imidazolate Frameworks Nanoparticles for Gene Delivery. *Microporous and Mesoporous Materials: The Official Journal of the International Zeolite Association* 302 (August): 110200. <https://doi.org/10.1016/j.micromeso.2020.110200>.
- Abdelhamid, Hani Nasser, Moataz Dowaidar, Mattias Hällbrink, and Ülo Langel. 2020. Gene Delivery Using Cell Penetrating Peptides-Zeolitic Imidazolate Frameworks. *Microporous and Mesoporous Materials: The Official Journal of the International Zeolite Association* 300 (June): 110173. <https://doi.org/10.1016/j.micromeso.2020.110173>.
- Ahmad, Almeman, Khalaf Hassan, Rasool Semaab, Moataz Dowaidar, and Al Orainy Mohammad. 2013. The Impact of CYP2C19 Polymorphism on Platelet Reactivity for Guiding Clopidogrel Treatment and Cost Analysis. *Journal of the Saudi Heart Association* 25 (2): 107. <https://doi.org/10.1016/j.jsha.2013.03.005>.
- Ai J, et al., 2008. Characterization of recombinant adeno-associated viral transduction and safety profiles in cardiomyocytes. *Cell. Physiol. Biochem* 48, 1894–1900.
- Akil O, Blits B, Lustig LR, Leake PA, 2019a. Virally mediated overexpression of glial-derived neurotrophic factor elicits age- and dose-dependent neuronal toxicity and hearing loss. *Hum. Gene Ther* 30 (1), 88–105.
- Akil O, Dyka F, Calvet C, Emptoz A, Lahlou G, Nouaille S, Boutet de Monvel J, Hardelin JP, Hauswirth WW, Avan P, Petit C, Safieddine S, Lustig LR, 2019b. Dual AAV-mediated gene therapy restores hearing in a DFNB9 mouse model. *Proc. Natl. Acad. Sci. U. S. A* 19.
- Akil O, Seal RP, Burke K, Wang C, Alemi A, During M, Edwards RH, Lustig LR, 2012. Restoration of hearing in the VGLUT3 knockout mouse using virally mediated gene therapy. *Neuron* 75 (2), 283–293. 10.1016/j.neuron.2012.05.019.
- Alam SA, Robinson BK, Huang J, Green SH, 2007. Prosurvival and proapoptotic intracellular signaling in rat spiral ganglion neurons in vivo after the loss of hair cells. *J. Comp. Neurol* 503 (6), 832–852.
- Alghasham, Abdullah, Ahmad A. A. Settin, Ahmad Ali, and Hisham Ismail. n.d. Association of MTHFR C677T and A1298C Polymorphisms with Hypertension among Saudi Subjects from Qassim Region. *International Journal of Health Sciences* 6 (1). Accessed June 18, 2021. <http://ijhs.org.sa/index.php/journal/article/view/312>.
- Alghasham, Abdullah, Hisham Ismail, Moataz Dowaidar, and Ahmad A. Settin. 2011. Methylenetetrahydrofolate Reductase (MTHFR) and Angiotensin Converting Enzyme (ACE) Gene Polymorphisms among Saudi Population from Qassim Region. *International Journal of Health Sciences* 5 (2 Suppl 1): 3–4. <https://www.ncbi.nlm.nih.gov/pubmed/23284552>.
- Alghasham, Abdullah, Ahmad A. Settin, Ahmad Ali, Moataz Dowaidar, and Hisham Ismail. 2012a. Association of MTHFR C677T and A1298C Gene Polymorphisms with Hypertension. *International Journal of Health Sciences* 6 (1): 3–11. <https://doi.org/10.12816/0005968>.
- Alghasham, Abdullah, Ahmad Ali, Hisham Ismail, Moataz Dowaidar, and Ahmad A. Settin. 2012. CYP2J2 -50 G/T and ADRB2 G46A Gene Polymorphisms in Saudi Subjects with Hypertension. *Genetic Testing and Molecular Biomarkers* 16 (9): 1027–31. <https://doi.org/10.1089/gtmb.2012.0006>.
- Ali, Ahmad, Abdullah Alghasham, Hisham Ismail, Moataz Dowaidar, and Ahmad Settin. 2013. ACE I/D and eNOS E298D Gene Polymorphisms in Saudi Subjects with Hypertension. *Journal of the Renin-Angiotensin-Aldosterone System: JRAAS* 14 (4): 348–53. <https://doi.org/10.1177/1470320312459976>.
- Ali, Ahmed A. A., Nahla M. Wassim, Moataz Dowaidar, and Ahmed E. Yaseen. 2013b. Association of eNOS (E298D) and CYP2J2 (-50G/T) Gene Polymorphisms with Hypertension among Egyptian Cases. *The Journal of Basic & Applied Zoology* 66 (4): 234–41. <https://doi.org/10.1016/j.jobaz.2012.12.001>.

- Ali, Ahmed A. A., Nahla M. Wassim, Moataz M. Dowaidar, and Ahmed E. Yaseen. 2013. Genetic Polymorphism of CYP2D6 Gene among Egyptian Hypertensive Cases. *The Journal of Basic & Applied Zoology* 66 (4): 228–33. <https://doi.org/10.1016/j.jobaz.2012.12.002>.
- Aljarallah, Badr, Ahmed Ali, Moataz Dowaidar, and Ahmad Settin. 2011. Prevalence of α -1-Antitrypsin Gene Mutations in Saudi Arabia. *Saudi Journal of Gastroenterology: Official Journal of the Saudi Gastroenterology Association* 17 (4): 256–60. <https://doi.org/10.4103/1319-3767.82580>.
- Alves S, Fol R, Cartier N, 2016. Gene therapy strategies for Alzheimer's disease: an overview. *Hum. Gene Ther* 27 (2), 100–107.
- Askew C, Rochat C, Pan B, Asai Y, Ahmed H, Child E, Schneider BL, Aebischer P, Holt JR, 2015. Tmc gene therapy restores auditory function in deaf mice. *Sci. Transl. Med* 7 (295), 295.
- Atkinson PJ, Wise AK, Flynn BO, Nayaga BA, Richardson RT, 2014a. Viability of long-term gene therapy in the cochlea. *Sci. Rep* 4, 4733.
- Atkinson PJ, Wise AK, Flynn BO, Nayagam BA, Hume CR, O'Leary SJ, Shepherd RK, Richardson RT, 2012. Neurotrophin gene therapy for sustained neural preservation after deafness. *PLoS One* 7 (12), e52338 [10.1371/journal.pone.0052338](https://doi.org/10.1371/journal.pone.0052338).
- Bainbridge JW, et al., Mehat MS, Sundaram V, Robbie SJ, Barker SE, Ripamonti C, 2015. Long-term effect of gene therapy on Leber's congenital amaurosis. *N Engl J Med* 372, 1887–1897. [10.1056/NEJMoa1414221](https://doi.org/10.1056/NEJMoa1414221). Epub 2015 May 4.
- Ballana E, Wang J, Venail F, Estivill X, Puel J-L, Arbones ML, Bosch A, 2008. Efficient and specific transduction of cochlear supporting cells by adeno-associated virus serotype 5. *Neurosci. Lett* 442 (2), 134–139. [10.1016/j.neulet.2008.06.060](https://doi.org/10.1016/j.neulet.2008.06.060).
- Bartus RT, Johnson EM Jr, 2017b. Clinical tests of neurotrophic factors for human neurodegenerative diseases, part 2: where do we stand and where must we go next? *Neurobiol. Dis* 97 (Pt B), 169–178.
- Bartus RT, Johnson EM Jr., 2017a. Clinical tests of neurotrophic factors for human neurodegenerative diseases, part 1: where have we been and what have we learned? *Neurobiol. Dis* 97 (Pt B), 156–168.
- Blesch A, Grill RJ, Tuszynski MH, 1998. Neurotrophin gene therapy in CNS models of trauma and degeneration. *Prog. Brain Res* 117, 473–484. Review.
- Blits B, Oudega M, Boer GJ, Bartlett Bunge M, Verhaagen J, 2003. Adeno-associated viral vector-mediated neurotrophin gene transfer in the injured adult rat spinal cord improves hind-limb function. *Neuroscience* 118 (1), 271–281.
- Boeda B, Weil D, Petit C, 2001. A specific promoter of the sensory cells of the inner ear defined by transgenesis. *Hum. Mol. Genet* 10, 1581–1589.
- Bowles DE, et al., McPhee SW, Li C, 2012. Phase 1 gene therapy for Duchenne muscular dystrophy using a translational optimized AAV vector. *Mol. Ther* 20, 443–455. [10.1038/mt.2011.237](https://doi.org/10.1038/mt.2011.237).
- Bringmann A, Pannicke T, Grosche J, Francke M, Wiedemann P, Skatchkov SN, Osborne NN, Reichenbach A, 2006. Müller cells in the healthy and diseased retina. *Prog. Retin. Eye Res* 25, 397–424.
- Budenz CL, Wong HT, Swiderski DL, Shibata SB, Pflug BE, Raphael Y, 2015. Differential effects of AAV.BDNF and AAV.Ntf3 in the deafened adult Guinea pig ear. *Sci. Rep* 5, 8619 [10.1038/srep08619](https://doi.org/10.1038/srep08619).
- Chang Q, Wang J, Li Q, Kim Y, Zhou B, Wang Y, Li H, Lin X, 2015. Virally mediated Kcnq1 gene replacement therapy in the immature scala media restores hearing in a mouse model of human Jervell and Lange-Nielsen deafness syndrome. *EMBO Mol. Med* 7 (8), 077–86.
- Chen H, Xing Y, Xia L, Chen Z, Yin S, Wang J, 2018. AAV-mediated NT-3 overexpression protects cochleae against noise-induced synaptopathy. *Gene Ther.* 25, 251–259. [10.1038/s41434-018-0012-0](https://doi.org/10.1038/s41434-018-0012-0).
- Chern A, Golub JS, 2019. Age-related hearing loss and dementia. *Alzheimer Dis. Assoc. Disord* [10.1097/WAD.0000000000000325](https://doi.org/10.1097/WAD.0000000000000325) ([Epub ahead of print]).
- Chien WW, Monzack EL, McDougald DS, Cunningham LL, 2015. Gene therapy for sensorineural hearing loss. *Ear Hear.* 36 (1), 1–7.
- Chikar JA, Colesa DJ, Swiderski DL, Di Polo A, Raphael Y, Pflug BE, 2008. Over-expression of BDNF by adenovirus with concurrent electrical stimulation improves cochlear implant thresholds and survival of auditory neurons. *Hear. Res* 245, 24–34. [10.1016/j.heares.2008.08.005](https://doi.org/10.1016/j.heares.2008.08.005).
- Cideciyan AV, et al., Jacobson SG, Beltran WA, Sumaroka A, Swider M, Iwabe S, 2013. Human retinal gene therapy for Leber congenital amaurosis shows advancing retinal degeneration despite enduring visual improvement. *Proc Natl Acad Sci USA* 110 (6), E517–E525. [10.1073/pnas.1218933110](https://doi.org/10.1073/pnas.1218933110). Epub 2013 Jan 22.

- Colella P, Ronzitti G, Mingozzi F, 2018. Emerging issues in AAV-mediated in vivo gene therapy. *Mol Ther Methods Clin Dev* 8, 87–104.
- Cunha C, Angelucci A, D'Antoni A, Dobrossy MD, Dunnett SB, Berardi N, Brambilla R, 2009. Brain-derived neurotrophic factor (BDNF) overexpression in the forebrain results in learning and memory impairments. *Neurobiol. Dis* 33 (3), 358–368.
- Daly C, Ward R, Reynolds AL, Galvin O, Collery RF, Kennedy BN, 2018. Brain-derived neurotrophic factor as a treatment option for retinal degeneration. *Adv. Exp. Med. Biol* 1074, 465–471. 10.1007/978-3-319-75402-4_57 (Review).
- Davis RL, 2003. Gradients of neurotrophins, ion channels, and tuning in the cochlea. *Neuroscientist* 9 (5), 311–316. Review.
- Di Polo A, Aigner LJ, Dunn RJ, Bray GM, Aguayo AJ, 1998. Prolonged delivery of brain-derived neurotrophic factor by adenovirus-infected Müller cells temporarily rescues injured retinal ganglion cells. *Proc. Natl. Acad. Sci. U. S. A* 95, 3978–3983.
- Dowaidar, M. Cell-Penetrating Peptides Uptake Pathways and Role in Drug Delivery with Potentials for Gene Therapy and Vaccine Development . Preprints.org 2023, 2023070889. <https://doi.org/10.20944/preprints202307.0889.v1>
- Dowaidar, M., J. Regberg, D. A. Dobchev, and T. Lehto. 2017. Refinement of a Quantitative Structure–activity Relationship Model for Prediction of Cell-Penetrating Peptide Based Transfection Systems. *International Journal of*. <https://link.springer.com/content/pdf/10.1007/s10989-016-9542-8.pdf>.
- Dowaidar, Moataz, and Ahmad Settin. 2010. Risk of Myocardial Infarction Related to Factor V Leiden Mutation: A Meta-Analysis. *Genetic Testing and Molecular Biomarkers* 14 (4): 493–98. <https://doi.org/10.1089/gtmb.2010.0017>.
- Dowaidar, Moataz, and Moataz Dowaidar. 2018. Chimeric Gene Delivery Vectors : Design, Synthesis, and Mechanisms from Transcriptomics Analysis.
- Dowaidar, Moataz, H. A. Ismail, A. A. Alghasham, M. M. Dowaidar, and A. A. Settin. 2011. Polymorphisms in MTHF and Ace Genes and the Association with Hypertension among Saudi Population from Qassim Region. *Egyptian Journal of Biochemistry and Molecular Biology* 29 (1). <https://doi.org/10.4314/ejbmb.v29i1.67382>.
- Dowaidar, Moataz, Hani Nasser Abdelhamid, Mattias Hällbrink, Krista Freimann, Kaido Kurrikoff, Xiaodong Zou, and Ülo Langel. 2017. Magnetic Nanoparticle Assisted Self-Assembly of Cell Penetrating Peptides-Oligonucleotides Complexes for Gene Delivery. *Scientific Reports* 7 (1): 9159. <https://doi.org/10.1038/s41598-017-09803-z>.
- Dowaidar, Moataz, Hani Nasser Abdelhamid, Mattias Hällbrink, Ülo Langel, and Xiaodong Zou. 2018. Supplemental Material for Chitosan Enhances Gene Delivery of Oligonucleotide Complexes with Magnetic Nanoparticles–cell-Penetrating Peptide. *SAGE Journals*. <https://doi.org/10.25384/SAGE.7105436.V1>.
- Dowaidar, Moataz, Hani Nasser Abdelhamid, Mattias Hällbrink, Ülo Langel, and Xiaodong Zou. 2018. Chitosan Enhances Gene Delivery of Oligonucleotide Complexes with Magnetic Nanoparticles-Cell-Penetrating Peptide. *Journal of Biomaterials Applications* 33 (3): 392–401. <https://doi.org/10.1177/0885328218796623>.
- Dowaidar, Moataz, Hani Nasser Abdelhamid, Mattias Hällbrink, Xiaodong Zou, and Ülo Langel. 2017. Graphene Oxide Nanosheets in Complex with Cell Penetrating Peptides for Oligonucleotides Delivery General Subjects. *Biochimica et Biophysica Acta, General Subjects*. <https://pubag.nal.usda.gov/catalog/5734174>.
- Dowaidar, Moataz. 2017. In-Silico Design of Peptide-Based Transfection Systems, in-Vitro Validation, and up-Take Pathways Investigation. Department of Neurochemistry, Stockholm University.
- Drennan WR, Rubinstein JT, 2008. Music perception in cochlear implant users and its relationship with psychophysical capabilities. *J. Rehabil. Res. Dev* 45, 779–789.
- Eaton MJ, Blits B, Ruitenber MJ, Verhaagen J, Oudega M, 2002. Amelioration of chronic neuropathic pain after partial nerve injury by adeno-associated viral (AAV) vector-mediated over-expression of BDNF in the rat spinal cord. *Gene Ther.* 9 (20), 1387–1395.
- Edwards TL, et al., Jolly JK, Groppe M, Barnard AR, Cottrell CL, Tolmachova T, 2016. Visual acuity after retinal gene therapy for choroideremia. *N Engl J Med.* 374, 1996–1998. 10.1056/NEJMc1509501. Epub 2016 Apr 21.
- Emptoz A, Michel V, Lelli A, Akil O, Boutet de Monvel J, Lahlou G, Meyer A, Dupont T, Nouaille S, Ey E, Franca de Barros F, Beraneck M, Dulon D, Hardelin JP, Lustig L, Avan P, Petit C, Safieddine S, 2017. Local gene therapy durably restores vestibular function in a mouse model of Usher syndrome type 1G. *Proc. Natl. Acad. Sci. U. S. A* 114 (36), 9695–9700.
- Fariñas I, Jones KR, Tessarollo L, Vigers AJ, Huang E, Kirstein M, de Caprona DC, Coppola V, Backus C, Reichardt LF, Fritsch B, 2001. Spatial shaping of cochlear innervation by temporally regulated neurotrophin expression. *J. Neurosci* 21, 6170–6180.
- Firszt JB, Holden LK, Skinner MW, Tobey EA, Petersen A, Gaggli W, Runge-Samuelson CL, Wackym PA, 2004. Recognition of speech presented at soft to loud levels by adult cochlear implant recipients of three cochlear implant systems. *Ear Hear.* 25, 375–387.

- Flores-Otero J, Xue HZ, Davis RL, 2007. Reciprocal regulation of presynaptic and postsynaptic proteins in bipolar spiral ganglion neurons by neurotrophins. *J. Neurosci* 19 27 (51), 14023–14034.
- Flotte TR, Trapnell BC, Humphries M, et al., 2011. Phase 2 clinical trial of a recombinant adeno-associated viral vector expressing alpha1-antitrypsin: interim results. *Hum. Gene Ther* 22, 1239–1247.
- Fritzsich B, Kersigo J, Yang T, Jahan I, Pan N, 2015. Neurotrophic factor function during ear development: expression changes define critical phases for neuronal viability In: Fay Richard R., Poper Arthur N. (Eds.), *The Spiral Ganglion Neurons*. Springer Handbook of Auditory Research, 2015. [Google Scholar]
- Fritzsich B, Pirvola U, Ylikoski J, 1999. Making and breaking the innervation of the ear: neurotrophic support during ear development and its clinical implications. *Cell Tissue Res*. 295, 369–382.
- Fritzsich B, Silos-Santiago I, Bianchi LM, Fariñas I, 1997. The role of neurotrophic factors in regulating the development of inner ear innervation. *Trends Neurosci*. 20 (4), 159–164 (Review).
- Fukui H, Wong HT, Beyer LA, Case BG, Swiderski DL, Di Polo A, Ryan AF, Raphael Y, 2012. BDNF gene therapy induces auditory nerve survival and fiber sprouting in deaf Pou4f3 mutant mice. *Sci. Rep* 2, 838 10.1038/srep00838.
- Gauthier R, Joly S, Pernet V, Lachapelle P, Di Polo A, 2005. Brain-derived neurotrophic factor gene delivery to muller glia preserves structure and function of light-damaged photoreceptors. *Invest. Ophthalmol. Vis. Sci* 46, 3383–3392.
- Géléoc GS, Holt JR, 2009. Sound strategies for hearing restoration. *Science* 344 (6184), 1241062.
- Geng R, Akil O, Gopal SR, Chen DH, Stepanyan R, Basch ML, Dinculescu A, Furness DN, Saperstein D, Hauswirth W, Lustig LR, Alagramam KN, 2017. Modeling and preventing progressive hearing loss in Usher syndrome III. *Sci. Rep* 7, 13480.
- Gestin, Maxime, Moataz Dowaidar, and Ülo Langel. 2017. Uptake Mechanism of Cell-Penetrating Peptides. *Advances in Experimental Medicine and Biology* 1030: 255–64. https://doi.org/10.1007/978-3-319-66095-0_11.
- Gillespie LN, Clark GM, Bartlett PF, Marzella PL, 2003. BDNF induced survival of auditory neurons in vivo: cessation of treatment leads to accelerated loss of survival effects. *J. Neurosci. Res* 71, 785–790. 10.1002/jnr.10542.
- Glueckert R, Bitsche M, Miller JM, Zhu Y, Prieskorn DM, Altschuler RA, Schrott-Fischer A, 2008. Deafferentation-associated changes in afferent and efferent processes in the Guinea pig cochlea and afferent regeneration with chronic intrascalar brain-derived neurotrophic factor and acidic fibroblast growth factor. *J. Comp. Neurol* 507, 1602–1621. 10.1002/cne.21619.
- György B, Sage C, Indzhukulian AA, Scheffer DI, Brisson AR, Tan S, Wu X, Volak A, Mu D, Tamvakologos PI, Li Y, Fitzpatrick Z, Ericsson M, Breakefield XO, Corey DP, Maguire CA, 2017. Rescue of hearing by gene delivery to inner-ear hair cells using exosome-associated AAV. *Mol. Ther* 25 (2), 379–391. 10.1016/j.ymthe.2016.12.010.
- Hardcastle N, Boulis NM, Federici T, 2018. AAV gene delivery to the spinal cord: serotypes, methods, candidate diseases, and clinical trials. *Expet Opin. Biol. Ther* 8 (3), 293–307.
- Harvey AR, Lovett SJ, Majda BT, Yoon JH, Wheeler LP, Hodgetts SI, 2015. Neurotrophic factors for spinal cord repair: which, where, how and when to apply, and for what period of time? *Brain Res*. 619, 36–71.
- Hashimoto K, Hickman TT, Suzuki J, Ji L, Corfas G, Liberman MC, 2019. Protection from noise-induced cochlear synaptopathy by virally mediated overexpression of NT3. *Sci Rep* 9 (1), 15362 10.1038/s41598-019-51724-6.
- Hendriks WT, Ruitenber MJ, Blits B, Boer GJ, Verhaagen J, 2004. Viral vector-mediated gene transfer of neurotrophins to promote regeneration of the injured spinal cord. *Prog. Brain Res* 46, 451–476. Review.
- Herzog KH, von Bartheld CS, 1998. Contributions of the optic tectum and the retina as sources of brain-derived neurotrophic factor for retinal ganglion cells in the chick embryo. *J. Neurosci* 18, 2891–2906.
- Hinderer C, et al., 2018. Severe toxicity in nonhuman primates and piglets following high-dose intravenous administration of an adeno-associated virus vector expressing human SMN. *Hum. Gene Ther* 29, 285–298.
- Hodgetts SI, Harvey AR, 2017. Neurotrophic factors used to treat spinal cord injury. *Vitam. Horm* 104, 405–457. Review.
- Holden LK, Finley CC, Firszt JB, Holden TA, Brenner C, Potts LG, Gotter BD, Vanderhoof SS, Mispagel K, Heydebrand G, Skinner MW, 2013. Factors affecting open-set word recognition in adults with cochlear implants. *Ear Hear*. 10.1097/AUD.0b013e3182741aa7.
- Huang EJ, Reichardt LF, 2001. Neurotrophins: roles in neuronal development and function. *Annu. Rev. Neurosci* 24, 677–736. Review.
- Isgrig K, Shteamer JW, Belyantseva IA, Drummond MC, Fitzgerald TS, Vijayakumar S, Jones SM, Griffith AJ, Friedman TB, Cunningham LL, Chien WW, 2017. Gene therapy restores balance and auditory functions in a mouse model of Usher syndrome. *Mol. Ther* 25 (3), 780–791.

- Ismail, H. A., A. A. Alghasham, M. M. Dowaidar, and A. A. Settin. 2011. Polymorphisms in MTHF and Ace Genes and the Association with Hypertension among Saudi Population from Qassim Region. *Egyptian Journal of Biochemistry and Molecular Biology* 29 (1). <https://doi.org/10.4314/ejbmb.v29i1.67382>.
- Jayakody DMP, Almeida OP, Speelman CP, Bennett RJ, Moyle TC, Yiannos JM, Friedland PL, 2018. Association between speech and high-frequency hearing loss and depression, anxiety and stress in older adults. *Maturitas* 110, 86–91. 10.1016/j.maturitas.2018.02.002.
- Jiam NT, Deroche ML, Jiradejvong P, Limb CJ, 2019. A randomized controlled crossover study of the impact of online music training on pitch and timbre perception in cochlear implant users. *J Assoc Res Otolaryngol* 20 (3), 247–262. 10.1007/s10162-018-00704-0.
- Jiao SS, Shen LL, Zhu C, Bu XL, Liu YH, Liu CH, Yao XQ, Zhang LL, Zhou HD, Walker DG, Tan J, Götz J, Zhou XF, Wang YJ, 2016. Brain-derived neurotrophic factor protects against tau-related neurodegeneration of Alzheimer's disease. *Transl. Psychiatry* 6 (10), e907.
- Johnson Chacko L, Blumer MJF, Pechriggl E, Rask-Andersen H, Dietl W, Haim A, Fritsch H, Glueckert R, Dudas J, Schrott-Fischer A, 2017. Role of BDNF and neurotrophic receptors in human inner ear development. *Cell Tissue Res.* 370 (3), 347–363. 10.1007/s00441-017-2686-9.
- Kamakura T, Nadol JB Jr., 2016. Correlation between word recognition score and intracochlear new bone and fibrous tissue after cochlear implantation in the human. *Hear. Res* 339, 132–141. 10.1016/j.heares.2016.06.015.
- Kanzaki S, Stöver T, Kawamoto K, Prieskorn DM, Altschuler RA, Miller JM, Raphael Y, 2002. Glial cell line-derived neurotrophic factor and chronic electrical stimulation prevent VIII cranial nerve degeneration following denervation. *J. Comp. Neurol* 454, 350–360. 10.1002/cne.10480.
- Keithley EM, Ma CL, Ryan AF, Louis JC, Magal E, 1998. GDNF protects the cochlea against noise damage. *Neuroreport* 9 (10), 2183–2187. 10.1097/00001756-199807130-00007.
- Khabou H, Cordeau C, Pacot L, Fisson S, Dalkara D, 2018. Dosage thresholds and influence of transgene cassette in adeno-associated virus-related toxicity. *Hum. Gene Ther* 29, 1235–1241.
- Khalin I, Alyautdin R, Kocherga G, Bakar MA, 2015. Targeted delivery of brain-derived neurotrophic factor for the treatment of blindness and deafness. *Int. J. Nanomed* 10, 3245–3267. 10.2147/IJN.S77480.eCollection. Review.
- Kilpatrick LA, Li Q, Goddard JC, Fekete DM, Lang H, 2011. Gene transfer using AVV viruses via the scala media in the adult normal and deafened mouse. *Gene Ther.* 18, 569–578.
- Klein RL, et al., 2006. Efficient neuronal gene transfer with AAV8 leads to neurotoxic levels of tau or green fluorescent proteins. *Mol. Ther* 13, 517–527.
- Koda M, Hashimoto M, Murakami M, Yoshinaga K, Ikeda O, Yamazaki M, Koshizuka S, Kamada T, Moriya H, Shirasawa H, Sakao S, Ino H, 2004. Adenovirus vector-mediated in vivo gene transfer of brain-derived neurotrophic factor (BDNF) promotes rubrospinal axonal regeneration and functional recovery after complete transection of the adult rat spinal cord. *J. Neurotrauma* 21 (3), 329–337.
- Kohrman DC, Wan G, Cassinotti L, Corfas G, 2019. Hidden hearing loss: a Disorder with multiple etiologies and mechanisms. *Cold Spring Harb Perspect Med* pii: a035493 [Epub ahead of print].
- Konishi M, Kawamoto K, Izumikawa M, Kuriyama H, Yamashita T, 2008. Gene transfer into Guinea pig cochlea using adeno-associated virus vectors. *J. Gene Med* 10 (6), 610–618. 10.1002/jgm.1189.
- Kovalchuk Y, Holthoff K, Konnerth A, 2004. Neurotrophin action on a rapid timescale. *Curr. Opin. Neurobiol* 14 (5), 558–563. Review.
- Kujawa SG, Liberman MC, 2009. Adding insult to injury: cochlear nerve degeneration after “temporary” noise-induced hearing loss. *J. Neurosci* 29, 14077–14085.
- Kujawa SG, Liberman MC, 2015. Synaptopathy in the noise-exposed and aging cochlea: primary neural degeneration in acquired sensorineural hearing loss. *Hear. Res* 330 (Pt B), 191–199. 10.1016/j.heares.2015.02.009.
- Landry TG, Fallon JB, Wise AK, Shepherd RK, 2013. Chronic neurotrophin delivery promotes ectopic neurite growth from the spiral ganglion of deafened cochleae without compromising the spatial selectivity of cochlear implants. *J. Comp. Neurol* 521 (12), 2818–2832. 10.1002/cne.23318.
- Lang H, 2015b. Loss, degeneration, and preservation In: Fay Richard R., Poper Arthur N. (Eds.), *The Spiral Ganglion Neurons*. Springer Handbook of Auditory Research. [Google Scholar]

- Lang H, Kilpatrick LA, Samuvel DJ, Krug EL, Goddard JC, 2011. Sox2 up-regulation and glial cell proliferation following degeneration of spiral ganglion neurons in adult mouse inner ear. *J. Assoc. Res. Otolaryngol* 12, 151–171.
- Lang H, Xing Y, Samuel DJ, Brown LN, Panganiban CH, Havens LT, Wegner M, Krug EL, Barth JL, 2015a. Neural stem/progenitor cell properties of glial cells in the adult mouse auditory nerve. *Sci. Rep* 5, 13383.
- Le Meur G, Lebranchu P, Billaud F, Adjali O, Schmitt S, Bezieau S, Pereon Y, Valabregue R, Ivan C, Darmon C, et al., 2018. Safety and long-term efficacy of AAV4 gene therapy in patients with RPE65 Leber congenital amaurosis. *Mol. Ther* 26, 256–268.
- Le TN, Straatman LV, Lea J, Westerberg B, 2017. Current insights in noise-induced hearing loss: a literature review of the underlying mechanism, pathophysiology, asymmetry, and management options. *J Otolaryngol Head Neck Surg* 46 (1), 41 10.1186/s40463-017-0219-x. Published 2017 May 23.
- Leake PA, Hradek GT, 1988. Cochlear pathology of long term neomycin induced deafness in cats. *Hear. Res* 33, 11–33.
- Leake PA, Hradek GT, Hetherington AM, Stakhovskaya O, 2011. Brain-derived neurotrophic factor promotes cochlear spiral ganglion cell survival and function in deafened, developing cats. *J. Comp. Neurol* 519, 1526–1545. 10.1002/cne.22582.
- Leake PA, Rebscher SJ, Dore 'C, Akil O, 2019. AAV-mediated neurotrophin gene therapy promotes improved survival of cochlear spiral ganglion neurons in neonatally deafened cats: comparison of AAV2-hBDNF and AAV5-hGDNF. *J Assoc Res Otolaryngol* 20 (4), 341–361. 10.1007/s10162-019-00723-5.
- Leake PA, Stakhovskaya O, Hetherington A, Rebscher SJ, Bonham B, 2013. Effects of brain-derived neurotrophic factor (BDNF) and electrical stimulation on survival and function of cochlear spiral ganglion neurons in deafened, developing cats. *J Assoc Res Otolaryngol* 14 (2), 187–211. 10.1007/s10162-013-0372-5.
- Lee CS, Bishop ES, Zhang R, Yu X, Farina EM, Yan S, Zhao C, Zheng Z, Shu Y, Wu X, et al., 2017. Adenovirus-mediated gene delivery: potential applications for gene and cell-based therapies in the new era of personalized medicine. *Genes Dis* 4, 43–63.
- Lieberman MC, 2015. Hidden hearing loss. *Sci. Am* 313, 48–53.
- Lieberman MC, 2017. Noise-induced and age-related hearing loss: new perspectives and potential therapies. *F1000Research* 6, 927 10.12688/f1000research.11310.1.
- Lieberman MC, Kujawa SG, 2017. Cochlear synaptopathy in acquired sensorineural hearing loss: manifestations and mechanisms. *Hear. Res* 349, 138–147. 10.1016/j.heares.2017.01.003.
- Liu Y, Okada T, Nomoto T, Ke X, Kume A, Ozawa K, Xiao S, 2007. Promoter effects of adeno-associated viral vector for transgene expression in the cochlea in vivo. *Exp. Mol. Med* 39, 170–175.
- Loera-Valencia R, Cedazo-Minguez A, Kenigsberg P, Page G, Duarte A, Giusti P, Zusso M, Robert P, Frisoni GB, Cattaneo A, Zille M, Boltze J, Cartier N, Buee L, Johansson G, Winblad B, 2019. Current and emerging avenues for Alzheimer's disease drug targets. *J. Intern. Med* 10.1111/joim.12959 ([Epub ahead of print] Review).
- Lustig LR, Akil O, 2012. Cochlear gene therapy. *Curr. Opin. Neurol* 25 (1), 57–60 submitted for publication.
- Ma Y, Wise AK, Shepherd RK, Richardson RT, 2019. New molecular therapies for the treatment of hearing loss. *Pharmacol. Ther* 200, 190–209. 10.1016/j.pharmthera.2019.05.003.
- MacLaren RE, et al., Groppe M, Barnard AR, Cottrill CL, Tolmachova T, Seymour L, 2014. Retinal gene therapy in patients with choroideremia: initial findings from a phase 1/2 clinical trial. *Lancet* 383, 1129–1137. 10.1016/S0140-6736(13)62117-0. Epub 2014 Jan 16.
- Malkki H, 2015. Alzheimer disease: NGF gene therapy activates neurons in the AD patient brain. *Nat. Rev. Neurol* 1 (10), 548.
- Mancino R, Martucci A, Cesareo M, Giannini C, Corasaniti MT, Bagetta G, Nucci C, 2018. Glaucoma and alzheimer disease: one age-related neurodegenerative disease of the brain. *Curr. Neuropharmacol* 167, 971–977. 10.2174/1570159X16666171206144045. Review.
- Martínez-Gálvez G, Zambrano JM, Diaz Soto JC, Zhan WZ, Gransee HM, Sieck GC, Mantilla CB, 2016. TrkB gene therapy by adeno-associated virus enhances recovery after cervical spinal cord injury. *Exp. Neurol* 276, 31–40.
- Maruyama J, Miller JM, Ulfendahl M, 2008. Glial cell line-derived neurotrophic factor and antioxidants preserve the electrical responsiveness of the spiral ganglion neurons after experimentally induced deafness. *Neurobiol. Dis* 29, 14–21. 10.1016/j.nbd.2007.07.026.

- McGee Sanftner LH, Abel H, Hauswirth WW, Flannery JG, 2001. Glial cell line derived neurotrophic factor delays photoreceptor degeneration in a transgenic rat model of retinitis pigmentosa. *Mol. Ther* 4, 622–629.
- Merentie M, et al., 2016. Efficacy and safety of myocardial gene transfer of adenovirus, adeno-associated virus and lentivirus vectors in the mouse heart. *Gene Ther.* 23, 296–305.
- Mestre TA, Sampaio C, 2017. Huntington Disease: linking pathogenesis to the development of experimental therapeutics. *Curr. Neurol. Neurosci. Rep* 17 (2), 18.
- Miller JM, Le Prell CG, Prieskorn DM, et al., 2007. Delayed neurotrophin treatment following deafness rescues spiral ganglion cells from death and promotes regrowth of auditory nerve peripheral processes: effects of brain-derived neurotrophic factor and fibroblast growth factor. *J. Neurosci. Res* 85, 1959–1969. 10.1002/jnr.21320.
- Mingozzi F, High KA, 2013. Immune responses to AAV vectors: overcoming barriers to successful gene therapy. *Blood* 122, 23–36.
- Moataz Dowaidar. 2012b. Association of MTHFR C677T and A1298C Gene Polymorphisms with Hypertension. *International Journal of Health Sciences* 6 (1): 3–11. <https://doi.org/10.12816/0005968>.
- Moataz Dowaidar. 2013. Association of eNOS (E298D) and CYP2J2 (–50G/T) Gene Polymorphisms with Hypertension among Egyptian Cases. *The Journal of Basic & Applied Zoology* 66 (4): 234–41. <https://doi.org/10.1016/j.jobaz.2012.12.001>.
- Moataz Dowaidar. 2017. Graphene Oxide Nanosheets in Complex with Cell Penetrating Peptides for Oligonucleotides Delivery. *Biochimica et Biophysica Acta, General Subjects* 1861 (9): 2334–41. <https://doi.org/10.1016/j.bbagen.2017.07.002>.
- Moataz Dowaidar. 2018. Chimeric Gene Delivery Vectors : Design, Synthesis, and Mechanisms from Transcriptomics Analysis. Department of Biochemistry and Biophysics, Stockholm University. <https://www.diva-portal.org/smash/record.jsf?pid=diva2:1242000>.
- Moataz Dowaidar. 2021gr. Exosomes Potential Therapeutics. <https://doi.org/10.31219/osf.io/mhwt3>.
- Moataz Dowaidar. 2D MOFs Have Unique Features for Biological Applications. They Can Be Utilized for Gene Therapy, Bioimaging, Biosensing, Photodynamic Therapy, and Tissue Engineering. <https://doi.org/10.31219/osf.io/4q9ct>.
- Moataz Dowaidar. 3D Bioprinting for Enhanced Vascularization, and Gene Editing to Provide a More Favorable Immunological Response Are Just Some of the Potential Uses of Carbon Materials. <https://doi.org/10.31219/osf.io/v2xy8>.
- Moataz Dowaidar. AAV9 Is Considered the Most Efficient AAV Serotype Targeting Blood-Brain Barriers. To Enhance Effective Gene Therapy for CNS Illnesses, Testing Novel Vectors with More Efficient Crossing Capabilities Is Vital. <https://doi.org/10.31219/osf.io/7bf5s>.
- Moataz Dowaidar. Addiction Biology Research on miRNAs, and Their Role in the Pathophysiology of Addiction Is Enabling Gene Therapy Opportunities. <https://doi.org/10.31219/osf.io/z5wyt>.
- Moataz Dowaidar. Anderson–Fabry Disease Can Be a Target for Gene Therapy. <https://doi.org/10.31219/osf.io/tcgka>.
- Moataz Dowaidar. Antisense Oligonucleotides (ASOs) and CRISPR Systems Are Promising Gene Therapy Treatments for Alzheimer’s Disease. <https://doi.org/10.31219/osf.io/ws796>.
- Moataz Dowaidar. Anti-Sense Pathways Have Been Generated Using siRNA. The Liver and Other Often Used Organs Will Now Be Targeted. <https://doi.org/10.31219/osf.io/m6xvp>.
- Moataz Dowaidar. Any Alteration in PPAR Genomic Sequence, Splicing Pattern, or PTM Is Likely to Cause Major Alterations in Its Function. In Personalized Medicine, Such Data Becomes More Significant in Gene Therapy Design. <https://doi.org/10.31219/osf.io/y8n79>.
- Moataz Dowaidar. Applying Genome-Wide Association Technology to Brain Diseases Enables the Discovery of lncRNAs Targets for Gene Therapy. <https://doi.org/10.31219/osf.io/hm4eu>.
- Moataz Dowaidar. Aptamers Targeting Vascular Endothelial Growth Factor Molecular Regulation as Potential Therapists. <https://doi.org/10.31219/osf.io/a8qpr>.
- Moataz Dowaidar. Arrhythmogenic Cardiomyopathy Is a Set of Hereditary Cardiac Muscle Disorders Where Various Etiologies Converge. Most ACM Patients Do Not Have a Genetic Diagnosis. <https://doi.org/10.31219/osf.io/pztv3>.
- Moataz Dowaidar. Artificial miRNAs Are Potential Gene Therapy Tools, Especially for Incurable Monogenic Disorders. <https://doi.org/10.31219/osf.io/d5rnm>.
- Moataz Dowaidar. Autophagy and Proteostasis Adjustment Role in Normal Brain Function and Neurodegenerative Disorders. <https://doi.org/10.31219/osf.io/m4yra>.
- Moataz Dowaidar. Autophagy, Immunological Response, and Inflammation All Rely on the TRIM Family Proteins. TRIM-Based Therapeutics for Inflammatory Illnesses Including Diabetes and Diabetic Comorbidities Are Promising. <https://doi.org/10.31219/osf.io/y4g6e>.
- Moataz Dowaidar. Bacterial Nanoparticles Can Deliver Proteins, Medications, Enzymes, and Genes to Diagnose and Cure Numerous Illnesses. <https://doi.org/10.31219/osf.io/7gyna>.
- Moataz Dowaidar. Basal Ganglia-Cerebellar and Brainstem-Cerebellar Circuits May Interact Improperly with Dystonia. Linking Network Disruptions to Cell Failure Will Enable Understanding Pathophysiology and Designing Gene Therapy Methods. <https://doi.org/10.31219/osf.io/8w35s>.

- Moataz Dowaidar. Biogenic Particles Can Be Multiantigenic, Immunostimulative and Activate Innate Immunity While Suppressing Tumor Development. <https://doi.org/10.31219/osf.io/q2kby>.
- Moataz Dowaidar. Biological Medications for Interventional Pain Have a Lot of Clinical Data behind Them. It Is Fair to Assume They Will Replace Steroid-Based Interventional Techniques, Providing Patients with Longer Relief. <https://doi.org/10.31219/osf.io/4y5fm>.
- Moataz Dowaidar. Blood Products Are Used to Treat a Multitude of Diseases, so the Blood Transfusion System Needs to Be Enhanced. CRISPR/Cas9 Has Made It Viable to Make HLA Class I-Deleted Blood Products to Avoid Rejection. <https://doi.org/10.31219/osf.io/egr3n>.
- Moataz Dowaidar. Breakthroughs in mRNA Modification and Nanoparticle-Based Delivery Vehicles Facilitate Gene Therapy Strategies. <https://doi.org/10.31219/osf.io/ky7dt>.
- Moataz Dowaidar. Calixarenes (CAs) Are Promising in Biomedicine, Biosensing, Bioimaging and Gene Delivery Systems. <https://doi.org/10.31219/osf.io/n9vjy>.
- Moataz Dowaidar. CAR T Cell Research Has Quickly Advanced from the Bench to the Clinic and Back. The Results of the Trials Have Revealed New Mechanisms. <https://doi.org/10.31219/osf.io/f9wm7>.
- Moataz Dowaidar. CAR T-Cell Treatment Remains Clinically Challenging. Therapeutic Strategies May Be Designed to Cut off Immunotherapy Utilizing Safety Switches. <https://doi.org/10.31219/osf.io/s7x4y>.
- Moataz Dowaidar. Carbon Nanofibers Assist in the Manufacture of Prosthetic Joints, Promote Tissue, Organ, Nerve Regeneration and Development, and Improve Anticancer Therapy Impact and Chemosensitization for a Range of Tumor Types. <https://doi.org/10.31219/osf.io/z3ucn>.
- Moataz Dowaidar. Carbon Nanotubes Have Enormous Potential in Gene Therapy. <https://doi.org/10.31219/osf.io/9bcxk>.
- Moataz Dowaidar. Cardiometabolic Conditions Could Be Related to Vitamin D Deficiency. The Genetic Determinants That Affect Vitamin D Pathways May Be Solved with Nanomedicines. <https://doi.org/10.31219/osf.io/nqewr>.
- Moataz Dowaidar. Central Nervous System Gene Therapy Has Entered a New Development Paradigm. New Techniques Are Being Employed for a Wide Range of Illness Indications and Pathways. <https://doi.org/10.31219/osf.io/j49wz>.
- Moataz Dowaidar. Charge-Alteration-Based Approaches Can Address the Evolving Needs of Nucleic Acid-Based Gene Therapy, Charge Reversal Techniques Are Also Promising. <https://doi.org/10.31219/osf.io/zwq5h>.
- Moataz Dowaidar. Chromosome X, the Most Explored Genome-Editing Chromosome, Presents Possibilities for Hemophilia A Treatments. <https://doi.org/10.31219/osf.io/6vsdz>.
- Moataz Dowaidar. Chronic Obstructive Pulmonary Condition (COPD) Is a Prevalent, Preventable, and Curable Illness with Persistent Respiratory Symptoms and Airflow Limitation. <https://doi.org/10.31219/osf.io/vkdut>.
- Moataz Dowaidar. CircRNAs Have the Potential to Aid in the Diagnosis and Treatment of Lipid Diseases. <https://doi.org/10.31219/osf.io/y3hp4>.
- Moataz Dowaidar. Clinical Investigations Show That siRNA May Be Used to Treat a Variety of Disorders, Including Cancer. <https://doi.org/10.31219/osf.io/fcsgq>.
- Moataz Dowaidar. Clinical Symptoms, Underlying Pathogenesis, and the Prospect of Tailored Therapies Have All Benefited from Genetic Discoveries in Parkinson's Disease. <https://doi.org/10.31219/osf.io/pdzqb>.
- Moataz Dowaidar. Code Distribution of siRNA for Cancer Genes such as p53 and Bcl2 Family Genes Has Demonstrated Efficacy in Killing Cancer Cells. Nanoparticles Can Produce a Surface Where Numerous Drugs May Be Coupled, Allowing Combinatory Treatment. <https://doi.org/10.31219/osf.io/hvcese>.
- Moataz Dowaidar. Cognitive Deficiencies Pathophysiology Are Mainly an Unknown Area. Curing the Neurological Conditions Could Be an Objective for Gene Therapy. <https://doi.org/10.31219/osf.io/23xf8>.
- Moataz Dowaidar. CRISPR/Cas System Research Has Advanced Significantly in Biological sciences. There Are Still Many Challenges to Effective Delivery before Efficient Gene Editing May Be Achieved. <https://doi.org/10.31219/osf.io/mc26v>.
- Moataz Dowaidar. CRISPR/Cas9 Genome Editing Technology Applications in Biological and Biomedical Fields. <https://doi.org/10.31219/osf.io/ctqbe>.
- Moataz Dowaidar. CRISPR/Cas9 Has Introduced New Gene Therapy Possibilities for Muscular Dystrophies. <https://doi.org/10.31219/osf.io/ug8v4>.
- Moataz Dowaidar. CRISPR/Cas9-Mediated Genome Editing Has Demonstrated Significant Promise for Genetic Correction in Autologous Hematopoietic Stem/progenitor Cells (HSPCs) and Induced Pluripotent Stem Cells (iPSCs). <https://doi.org/10.31219/osf.io/xk54r>.
- Moataz Dowaidar. CrisPR/CRIS Systems Are Highly Effective and Useful for Genomic Manipulation. Despite This, Cardiac Treatment Remains Difficult due to Existing Genome Editing and Delivery Processes. <https://doi.org/10.31219/osf.io/3nwzd>.
- Moataz Dowaidar. CRISPR-Based Gene Editing Is Presently Being Tried in Many Clinical Trials. <https://doi.org/10.31219/osf.io/qbngx>.
- Moataz Dowaidar. CRISPR-Cas9 Gene Editing as a Tool for Developing Immunotherapy for Cancer. <https://doi.org/10.31219/osf.io/dvr4t>.
- Moataz Dowaidar. Critical Limb Ischemia Potential Gene Therapy Strategies. <https://doi.org/10.31219/osf.io/aqcpt>.
- Moataz Dowaidar. Cyclodextrins as Potential Gene Therapy Vectors. <https://doi.org/10.31219/osf.io/zhtsc>.

- Moataz Dowaidar. Deep Learning Algorithms for scRNAseq Analysis Have Yielded Positive Results, but There Are Still More Promising Ways That Need to Be Developed for Regenerative Medicine. <https://doi.org/10.31219/osf.io/dh2pt>.
- Moataz Dowaidar. Degradable Branched Polycationic Systems Are Promising Gene Therapy Vectors. <https://doi.org/10.31219/osf.io/utypf>.
- Moataz Dowaidar. Depression May Be Epigenetically Controlled by miRNAs Making It a Diagnostic or Gene Therapy Target. <https://doi.org/10.31219/osf.io/fw65m>.
- Moataz Dowaidar. Dermatophytes: Role of Host Genetics in the Development of Illness. <https://doi.org/10.31219/osf.io/mf3bu>.
- Moataz Dowaidar. Developing Nanotechnology Platforms for Peptide-Based Combinatory Cancer Gene Therapy Will Likely Have a Significant Influence on the Development of Personalized Cancer Medicines. <https://doi.org/10.31219/osf.io/zbrkj>.
- Moataz Dowaidar. Development of Specialized Carriers Capable of Delivering Effective RNAi and siRNA Gene Therapy. <https://doi.org/10.31219/osf.io/3ykwk>.
- Moataz Dowaidar. Developments in Biomedical Technology Will Increase the Importance of mRNA in Treating Brain Tumors, as Well as Other Malignancies. <https://doi.org/10.31219/osf.io/tvj5x>.
- Moataz Dowaidar. Different Insulin Resistance and Inflammation Pathways Are Influenced by Genetic Factors in Metabolic Syndrome. Gene Therapy Enables Early Recognition and Treatment of the Genetic Factors. <https://doi.org/10.31219/osf.io/gqwj2>.
- Moataz Dowaidar. Discoveries in Gene-Environment Interactions That Influence CVD, Lipid Traits, Obesity, Diabetes, and Hypertension Appear to Be Able to Influence Gene Therapy. <https://doi.org/10.31219/osf.io/cr5af>.
- Moataz Dowaidar. Downstream Processing of Virus, Virus-like Particles and Nanoparticulate Inclusion Bodies to Be Used as Gene Delivery Vehicles for Human Gene Therapy Applications. <https://doi.org/10.31219/osf.io/exa3q>.
- Moataz Dowaidar. Dravet Syndrome Is a Severe Developmental and Epileptic Encephalopathy. Fenfluramine and Gene Therapy Are Promising. <https://doi.org/10.31219/osf.io/zvq8y>.
- Moataz Dowaidar. Emerging Therapy Options May Help Patients with RAG Deficiency, Especially Those with Severe Immune Dysregulation. <https://doi.org/10.31219/osf.io/v5tjg>.
- Moataz Dowaidar. Exosomal miRNA Diagnostic and Gene Therapy Tools. <https://doi.org/10.31219/osf.io/aknrc>.
- Moataz Dowaidar. Exosomes as Promising Gene Therapy Tools Still Need to Be Researched and Manufactured More Efficiently. <https://doi.org/10.31219/osf.io/nw4z7>.
- Moataz Dowaidar. Exosomes Can Make the Use of Circulating miRNA as a Biomarker More Feasible. The Aim of Gene Therapy Should Be to Learn Everything There Is to Know about miRNA Activity. <https://doi.org/10.31219/osf.io/edkua>.
- Moataz Dowaidar. Exosomes May Prevent Cardiac Attacks, Heart Failure, and Cardiomyopathy. <https://doi.org/10.31219/osf.io/agm3k>.
- Moataz Dowaidar. Exosomes' Function in Cardiovascular Protection and Neovascularization Implies That They Might Be Used to Treat Ischemia and Atherosclerotic Cardiovascular Diseases. <https://doi.org/10.31219/osf.io/2h8c7>.
- Moataz Dowaidar. Ferropoptosis Cell Death Can Cause Complications That May Be Difficult to Detect and Quantify: Autophagy Role and Possible Therapeutics. <https://doi.org/10.31219/osf.io/zd2jg>.
- Moataz Dowaidar. Focus on Exosomes Could Help Make the Use of Circulating miRNA as Biomarkers More Practical. A Detailed Understanding of miRNA Behavior Should Be a Subject of Gene Therapy. <https://doi.org/10.31219/osf.io/uan6x>.
- Moataz Dowaidar. Following the Discovery of Anti-MDA5 Ab, the Clinical Understanding of Dermatomyositis Has Been Improved. <https://doi.org/10.31219/osf.io/j2t5f>.
- Moataz Dowaidar. For the Treatment of Cystic Fibrosis, RNA Medicines, Gene Transfer Therapies, and Gene Editing Treatments Have Potential. <https://doi.org/10.31219/osf.io/6afzm>.
- Moataz Dowaidar. Frontotemporal Dementia Is a Complex Disorder with a Wide Spectrum of Clinical Symptoms. Personalized Medicine and Gene Therapy Are Promising Strategies for Treatment. <https://doi.org/10.31219/osf.io/gh4x7>.
- Moataz Dowaidar. G6PD Deficiency Is a Common Genetic Trait That Can Protect Heterozygotes from Dying from Malaria. <https://doi.org/10.31219/osf.io/g2kza>.
- Moataz Dowaidar. Gastric Cancer Is the World's Second-Largest Death Cause. Peptides Can Be Used to Deliver Radiation or Other Fatal Chemicals to Tumors. <https://doi.org/10.31219/osf.io/eu5mj>.
- Moataz Dowaidar. Gene Doping May Be Possible for Lifestyle Enhancement. <https://doi.org/10.31219/osf.io/8xkm5>.
- Moataz Dowaidar. Gene Expression Assays Gather Evidence That They Can Provide Useful Therapeutic Information in Young Women. <https://doi.org/10.31219/osf.io/d372s>.
- Moataz Dowaidar. Gene Modification Research Has Potential, from Diagnostic to Therapeutic Levels. The Most Promising Metabolic Pathways Include the TGF-1 Signaling System, Inflammation and Protein Transport. <https://doi.org/10.31219/osf.io/5ert4>.
- Moataz Dowaidar. Gene Therapy and Genome-Editing Treatments That Can Protect Patients from Coronary Artery Disease Are under Investigation. <https://doi.org/10.31219/osf.io/xqgf8>.
- Moataz Dowaidar. Gene Therapy Approaches for Hemophilia A and B. <https://doi.org/10.31219/osf.io/ufc4g>.

Moataz Dowaidar. Gene Therapy Can Target Mutations such as BRAF, Which Have Been Shown to Make Tumors More Susceptible to Autophagy Suppression. <https://doi.org/10.31219/osf.io/3gwra>.

Moataz Dowaidar. Gene Therapy Development and Legislation. <https://doi.org/10.31219/osf.io/mwb2n>.

Moataz Dowaidar. Gene Therapy for the Central Nervous System Has Been Initiated. This Expansion Will Require Some Degree of Simplicity in Delivery Processes. <https://doi.org/10.31219/osf.io/hdy5q>.

Moataz Dowaidar. Gene Therapy for the Treatment of Spinal Muscular Atrophy. <https://doi.org/10.31219/osf.io/kpz5f>.

Moataz Dowaidar. Gene Therapy Has Been Shown to Be Valuable for Understanding Complex Disease Pathophysiologies. The Medical Profession as a Whole Will Have to Invest in Specialized Investigations. <https://doi.org/10.31219/osf.io/8fg9y>.

Moataz Dowaidar. Gene Therapy May Benefit Inherited Ichthyoses with Concurrent Fungal Infections and Severe Ichthyoidoses. <https://doi.org/10.31219/osf.io/zxmun>.

Moataz Dowaidar. Gene Therapy May Target APOE for Alzheimer's Disease. <https://doi.org/10.31219/osf.io/3y52k>.

Moataz Dowaidar. Gene Therapy Promises Accurate, Targeted Administration and Overcoming Drug Resistance in Diverse Cancer Cells. <https://doi.org/10.31219/osf.io/j34n6>.

Moataz Dowaidar. Gene Therapy Targeting FVIII, FIX for Haemophilia Treatment. <https://doi.org/10.31219/osf.io/qcbwp>.

Moataz Dowaidar. Gene Therapy Targeting PRMT5 May Be Useful in Immunotherapy. <https://doi.org/10.31219/osf.io/gkw8j>.

Moataz Dowaidar. Gene Therapy Using Extracellular Vesicles Loaded with miRNA Derived from Bone Marrow Mesenchymal Stem Cells Is a Cell-Free Medication Delivery Method Used in a Variety of Diseases. <https://doi.org/10.31219/osf.io/3znvw>.

Moataz Dowaidar. Gene Therapy Using miRNA Treatment Suppresses the Expression of Bone-Forming Defective Genes and Raises the Expression of Genes That Become Dormant during Bone Building. <https://doi.org/10.31219/osf.io/tcka3>.

Moataz Dowaidar. Gene Therapy Using MnO₂ Nanoparticles. <https://doi.org/10.31219/osf.io/xmwjs>.

Moataz Dowaidar. Gene Therapy Vectors for Targeting the Heart. <https://doi.org/10.31219/osf.io/gcbhf>.

Moataz Dowaidar. Gene Therapy Vectors Should Enable CRISPR Systems to Accumulate at Disease Sites and Successfully Penetrate Nuclei. <https://doi.org/10.31219/osf.io/xzmmc>.

Moataz Dowaidar. Gene-Free Viral-like Particles (VLPs) Offer a Safer Alternative to Inactivating or Weakening Viral Strains for Traditional Vaccines. VLP-Based Vaccinations without Adjuvants Have Been Found to Promote Humoral and Cellular Immunity. <https://doi.org/10.31219/osf.io/9dvut>.

Moataz Dowaidar. Gene-Regulatory Elements May Change the Amount, Timing, or Location of Gene Expression, Cis-Regulation Therapy Platforms Might Become a Gene Therapy to Treat Many Genetic Diseases. <https://doi.org/10.31219/osf.io/xc5a2>.

Moataz Dowaidar. Genetic and Epigenetic Discoveries Hold Promising Avenues in Cardiovascular Prevention and Management (CVDs). Key Nucleic Acids Are Being Researched and Developed for Medicinal Use. <https://doi.org/10.31219/osf.io/hk7pe>.

Moataz Dowaidar. Genetic Engineered MSCs Are Attractive Possibilities for Regenerative Stem-Cell Therapy to Treat Several Liver Diseases. <https://doi.org/10.31219/osf.io/4cfrd>.

Moataz Dowaidar. Genetic Variants Shared between Alzheimer's Disease and Parkinson's Disease Have Been Discovered in Blood and Brain Samples. Somatic Mosaicism Might Function as an Accelerator. <https://doi.org/10.31219/osf.io/tr58n>.

Moataz Dowaidar. Genome Editing Can Now Be Carried out in an Isogenic Setting. It Can Be Effectively Transmitted to Somatic Tissues in Mice, but Not to Humans. Despite These Doubts, CRIS Has Great Potential as a Medical Promise. <https://doi.org/10.31219/osf.io/4rn3v>.

Moataz Dowaidar. Genome Editing's Potential Target Diseases in the Cardiovascular Field. <https://doi.org/10.31219/osf.io/ge23p>.

Moataz Dowaidar. Genome-Editing Is Promising for Producing Therapeutically Relevant Animal Models for Possible Therapies for Rare Human Diseases. <https://doi.org/10.31219/osf.io/dehr9>.

Moataz Dowaidar. Genome-Wide Association Experiments Have Uncovered a Slew of Cardiometabolic Trait-Associated Variants. This Information Can Be Useful in the Implementation of New Diagnostic and Treatment Strategies. <https://doi.org/10.31219/osf.io/4vws8>.

Moataz Dowaidar. Genome-Wide Association Studies (GWAS) Have Revolutionized Our View of Human Health and Disease Genetics and Offered Novel Gene Therapy Targets. <https://doi.org/10.31219/osf.io/rvm3z>.

Moataz Dowaidar. Genome-Wide Association Studies Promise to Discover Novel Indicators of Hypertension. Endothelin-Related SNPs Are Currently in Clinical Trials. <https://doi.org/10.31219/osf.io/2n4wa>.

Moataz Dowaidar. Gingival and Intraventricular Haemorrhages Are Severe Newborn Diseases Causing Damage to White Matter and Neurological Dysfunction in Surviving Newborns Who Can Benefit from Gene Therapy. <https://doi.org/10.31219/osf.io/qb84p>.

Moataz Dowaidar. Given the Importance of mTOR Signaling in a Number of Illnesses, It Looks Suitable to Use miR 99 Family Members as a Therapeutic Intervention to Deal with These Illnesses by Using Gene Therapy Tools. <https://doi.org/10.31219/osf.io/8cwgh>.

Moataz Dowaidar. Glioblastoma Therapeutic Approaches Were Established Utilizing Contemporary Discoveries in Delivering Medicines to the Brain as Smart Nanoparticles for Focused Therapy. <https://doi.org/10.31219/osf.io/db4f6>.

Moataz Dowaidar. Haemophilia Gene Therapy Is in Clinical Studies, Making Continuous Safety and Efficacy Testing a Key Emphasis. <https://doi.org/10.31219/osf.io/sa8ny>.

Moataz Dowaidar. Hematopoietic Stem Cell Transplantation and Gene Therapy Are the Sole Treatments for Sickle Cell Disease and Other Hemoglobinopathies. <https://doi.org/10.31219/osf.io/v8xqc>.

Moataz Dowaidar. Hemophilia Therapeutics. <https://doi.org/10.31219/osf.io/gu74x>.

Moataz Dowaidar. HMGB1 Has Sparked a Lot of Attention as a Model DAMP Molecule Involved in Inflammation, Inflammatory Diseases, and Cancer. <https://doi.org/10.31219/osf.io/5qx36>.

Moataz Dowaidar. Human Corneal Endothelial Cells Grafts to Replace Cadaveric Donor Corneas. <https://doi.org/10.31219/osf.io/p9x7e>.

Moataz Dowaidar. Huntington's Disease Gene Therapy and Nanomedicines May Be Available Shortly. <https://doi.org/10.31219/osf.io/rxvgd>.

Moataz Dowaidar. Hybrid Gene Therapy Designed to Fully Understand the Underlying Molecular Cancer Process May Be a Feasible Option. <https://doi.org/10.31219/osf.io/ajyfd>.

Moataz Dowaidar. Hybrid Nanotechnology and Peptide Nucleic Acid Could Improve the Effectiveness of Gene Therapy by Increasing Its Cell Permeability. <https://doi.org/10.31219/osf.io/d8wzt>.

Moataz Dowaidar. Hydrogels Are Promising Considering Their Incredible Capacity to Modify, Encapsulate and Co-Deliver Medicinal Compounds, Cells, Biomolecules, and Nanomaterials. <https://doi.org/10.31219/osf.io/px3qy>.

Moataz Dowaidar. Immune Evasion Is Linked to Histone Variation Malfunction. Gene Therapy Could Provide Tools for Targeting Histone Variant Deposition as a Critical Part of Its Pharmacology. <https://doi.org/10.31219/osf.io/kjm76>.

Moataz Dowaidar. Implementing the Human Artificial Chromosome Gene Therapy Platform Remains Challenging, but Continuous Animal Model Research Will Advance the Platform Closer to Clinical Trials. <https://doi.org/10.31219/osf.io/a53f7>.

Moataz Dowaidar. In Prenatal Stem Cell Transplantation and in Utero Gene Therapy, a Wide Spectrum of Genetic Diseases Can Be Diagnosed and Treated before Birth. <https://doi.org/10.31219/osf.io/sa3vz>.

Moataz Dowaidar. Inflammatory Breast Cancer Remains the Most Aggressive Form of Breast Cancer. A Multimodality Therapeutic Plan Has Shown Improved Survival Results. <https://doi.org/10.31219/osf.io/cr935>.

Moataz Dowaidar. Inherited Immunohematological and Metabolic Diseases Have the Potential to Improve Significantly, or Be Cured, Using Haematopoietic Stem Cell Transplantation Gene Therapy. <https://doi.org/10.31219/osf.io/ukbnm>.

Moataz Dowaidar. Insulin and IGF-1 Receptors Mutations Can Lead to Targets for Gene Therapy in Diabetes, Obesity, and Metabolic Syndrome. <https://doi.org/10.31219/osf.io/s86x5>.

Moataz Dowaidar. Integrating High-Throughput Genetics and Neuroimaging Technologies Promises Greater Information on Neurobiological Anomalies in Neurodegenerative Diseases. <https://doi.org/10.31219/osf.io/hpgyz>.

Moataz Dowaidar. Intravitreal and Subretinal Injections Currently Deliver Most Gene Therapy, Including siRNA for Eye Illnesses. Non-Viral Vectors May Provide Targeting. <https://doi.org/10.31219/osf.io/rjkhy>.

Moataz Dowaidar. Key Genetic Factors in the Metabolic Syndrome Predisposition Which May Be a Therapeutic Options by Gene Therapy. <https://doi.org/10.31219/osf.io/f38sk>.

Moataz Dowaidar. Liposomes Can Minimize Cardiotoxicity, Address Drug Resistance, and Improve Overall Drug Release Profiles in Breast Cancer. <https://doi.org/10.31219/osf.io/tn56d>.

Moataz Dowaidar. Liposomes with Cerasome-Forming Lipids as Gene Therapy Vectors. <https://doi.org/10.31219/osf.io/zjn6v>.

Moataz Dowaidar. LncRNA Regulating Reprogramming Glucose Metabolism Has Become One of the Most Tempting Antineoplastic Targets for Gene Therapy. <https://doi.org/10.31219/osf.io/hqma5>.

Moataz Dowaidar. lncRNAs Are Upregulated and Downregulated in OS Cells. Angiogenesis, Metastasis, Cell Signaling, Autophagy, and Death Are among Biological Processes That RNAs Play a Role in. <https://doi.org/10.31219/osf.io/48n7q>.

Moataz Dowaidar. Magnetic Iron Oxide Nanoparticles Have Potential on Gene Therapy Effectiveness and Biocompatibility. <https://doi.org/10.31219/osf.io/f3hm4>.

Moataz Dowaidar. Magnetic Nanoparticles Are Widely Used in Drug Delivery, Imaging, Diagnosis, and Targeting. It Has Promises for the Treatment of Inflammatory Disorders such as Rheumatoid Arthritis. <https://doi.org/10.31219/osf.io/p2gme>.

Moataz Dowaidar. Many miRNAs Participate in Inflammatory Regulation and Bone Metabolism. Overexpression of miR21 and miR155 Releases Proinflammatory Cytokines. <https://doi.org/10.31219/osf.io/2wuvp>.

Moataz Dowaidar. Mesenchymal Stem Cells Strategies in Cancer Immunotherapy. <https://doi.org/10.31219/osf.io/dkv6w>.

Moataz Dowaidar. Metabolic Syndrome_ the Presence of Inflammatory Mechanisms in Abdominal Obesity Is Undeniable, Gene Therapy Using Nanoparticles and Adenoviruses Technologies Is Promising. <https://doi.org/10.31219/osf.io/2j5xt>.

Moataz Dowaidar. MiR490's Diagnostic Capacity Was Demonstrated in Various Cancer Kinds and Diseases, Adding to Its Clinical Value. <https://doi.org/10.31219/osf.io/wysre>.

Moataz Dowaidar. miRNA Can Be a Part of Both the Onset and Cure of Coronary Heart Disease. <https://doi.org/10.31219/osf.io/teqh8>.

- Moataz Dowaidar. miRNAs Have an Impact on Xeno-Infectious Diseases by Influencing Host And/or Infection Factors. <https://doi.org/10.31219/osf.io/7qewx>.
- Moataz Dowaidar. miRNAs May Be Used as Preventive Agents for Metabolic Diseases in the near Future. Understanding the Interplay between pro-Adipogenic_ and Anti-Ad Pipogenic miRNA' Could Lead to New Biomarkers. <https://doi.org/10.31219/osf.io/3dr8c>.
- Moataz Dowaidar. Mutations in MED12 Lead to Mental Retardation, Including Opitz–Kaveggia Syndrome, Ohdo Syndrome, Lujan–Fryns Syndrome, and Psychosis. It's a Target for Gene Therapy. <https://doi.org/10.31219/osf.io/cyns8>.
- Moataz Dowaidar. Nanocarriers Can Be Used to Control the Activity of Genome Editing in a Spatiotemporal Way by Using Stimulusresponsive Nanocarriers. <https://doi.org/10.31219/osf.io/nua89>.
- Moataz Dowaidar. Nanoformulations Can Be Utilized to Deliver Effective siRNA to Tumor Cells to Decrease Gene Expression. <https://doi.org/10.31219/osf.io/zvukc>.
- Moataz Dowaidar. Nanomaterials Can Inhibit Planktonic and Biofilm Bacteria and Can Be Used as Topical Therapy for Mouth and Wound-Related Infections. <https://doi.org/10.31219/osf.io/aqd2e>.
- Moataz Dowaidar. Nanomaterials Combine Multiple Therapeutic Approaches for Cancer Cell Multidrug Resistance, Ferroptotic Cell Death Is Promising in Various Cancers. <https://doi.org/10.31219/osf.io/7bg9t>.
- Moataz Dowaidar. Nanomaterials Were Formed into Various Shapes, with Functionalization Aimed at Various Internalization Processes. Their Nanoscale Size Allows Drugs to Reach Cells or Extracellular Environments. <https://doi.org/10.31219/osf.io/p2ajv>.
- Moataz Dowaidar. Nanomedicine Has Elegantly Attempted to Cure Multiple Gene Polymorphisms and Mutations in Cardiovascular Diseases Using Gene Therapy Techniques. <https://doi.org/10.31219/osf.io/d3x8g>.
- Moataz Dowaidar. Nanomedicine Is Offering Promising Strategies for Tumor Blockade Treatment. <https://doi.org/10.31219/osf.io/yzxuq>.
- Moataz Dowaidar. Nanomedicines for Enhanced Permeability and Retention (EPR)-Stratified Patients Have the Potential to Improve Treatment Outcomes. <https://doi.org/10.31219/osf.io/xrcb2>.
- Moataz Dowaidar. Network Medicine Might Lead to New Treatments for Dyslipidemia. It Will Be a Challenging Method to Implement in a Clinical Context. <https://doi.org/10.31219/osf.io/nksbw>.
- Moataz Dowaidar. Neuroinflammation Caused by Activated Microglia and Astrocytes Can Contribute to the Progression of Pathogenic Damage to Substantia Nigra Neurons, Playing a Role in Parkinson's Disease Progression. <https://doi.org/10.31219/osf.io/ac896>.
- Moataz Dowaidar. Neurologists Rarely Perform Genetic Testing for Parkinson's Disease. Evidence Suggests That Many Patients with Major Genetic Variants Go Undiagnosed. <https://doi.org/10.31219/osf.io/ykpb2>.
- Moataz Dowaidar. Neuronal Ceroid Lipofuscinosis Therapeutics. <https://doi.org/10.31219/osf.io/75vcp>.
- Moataz Dowaidar. Neuronal Intranuclear Hyaline Inclusion Disease Is a Neurodegenerative Condition Which Can Be a Target for Gene Therapy. <https://doi.org/10.31219/osf.io/upgqd>.
- Moataz Dowaidar. Neurotrophin Gene Therapy May Be Able to Treat Individuals with Noise-Induced Hearing Loss or Neural Presbycusis. <https://doi.org/10.31219/osf.io/spkxh>.
- Moataz Dowaidar. New Technologies to Improve CAR T Cell Generation and Biomanufacturing Will Lead to Safer, More Therapeutically Effective Cells. <https://doi.org/10.31219/osf.io/un8gp>.
- Moataz Dowaidar. New Therapies Aim at Restoring the Molecular, Morphological, and Functional Integrity of Parkinson's Specific Brain Circuits. <https://doi.org/10.31219/osf.io/dvyxc>.
- Moataz Dowaidar. Next-Generation Sequencing Is Now Utilized to Identify Genetic Abnormalities and Develop Gene Therapy. <https://doi.org/10.31219/osf.io/em7xp>.
- Moataz Dowaidar. Nonviral Gene Delivery Vectors for Transfection of the CAR Gene for CAR-T Cell Therapy. <https://doi.org/10.31219/osf.io/ckxh5>.
- Moataz Dowaidar. Not All lncMIRHG's Are 'Junk Transcripts,'. lncMIRHG Loci May Make Both Functional miRNAs and lncRNAs, Which Can Work Together or Separately. <https://doi.org/10.31219/osf.io/a567w>.
- Moataz Dowaidar. Nrf2 Signaling Pathways Are Part of a Wider Network of Signaling Pathways Regulating Thymoquinone Therapeutic Actions Which Need Innovative Formulations and Delivery Methods. <https://doi.org/10.31219/osf.io/u2fa7>.
- Moataz Dowaidar. Nucleic Acid Designs, Artificial Intelligence for Screening Nanomaterials, and Enhanced Characterization Methods Are Needed to Make Nanomedicine More Successful. <https://doi.org/10.31219/osf.io/2w5aq>.
- Moataz Dowaidar. Nucleic Acid Nanocarriers Can Be Programmable, Spatially Adjustable and Biocompatible, Minimizing Systemic Toxicity and Improving Pharmacodynamics. <https://doi.org/10.31219/osf.io/wr237>.
- Moataz Dowaidar. Ocular Gene Therapy Strategies. <https://doi.org/10.31219/osf.io/7en3k>.
- Moataz Dowaidar. Omics Should Be Integrated with Genomics to Uncover Molecular Networks and Tissue and Single-Cell Epigenetic Changes. With These Findings, Targeted Pseudoexfoliation Syndrome and Glaucoma Gene Therapy Procedures May Be Viable. <https://doi.org/10.31219/osf.io/48fj5>.
- Moataz Dowaidar. Ophthalmic Gene and Cell Therapies. <https://doi.org/10.31219/osf.io/n84m9>.

Moataz Dowaidar. Osteoporosis Is a Prominent Source of Morbidity and Mortality in the Elderly, Particularly in Postmenopausal Women. Long Noncoding RNAs (lncRNAs) Have Been Found to Be Important Regulators and Possible Gene Therapy Targets. <https://doi.org/10.31219/osf.io/ghfpt>.

Moataz Dowaidar. p21 Is a Flexible, Multi-Functional Protein. It Governs Various Tumor Cell Activities, Including Autophagy. p21 Is a Possible Radiotherapy Target. <https://doi.org/10.31219/osf.io/ydkca>.

Moataz Dowaidar. Parkinson's Disease Simulating Complexity via Improving the Identification of Significant Genetic Alterations and Environmental Contaminants Should Be a Priority. <https://doi.org/10.31219/osf.io/pmcu9>.

Moataz Dowaidar. Patients with PMD Who Are Thoroughly Screened by Genomic Medicine Have a Considerable Chance of Benefiting Greatly from Whole-Genome Sequencing. <https://doi.org/10.31219/osf.io/dajft>.

Moataz Dowaidar. Patient-Specific Microphysiology Systems Are Likely to Become a Crucial Aspect of Translational Research and Precision Medicine. <https://doi.org/10.31219/osf.io/bc8fr>.

Moataz Dowaidar. Peripheral Nerve Injury Therapeutics, Including Electrical Stimulation, Stem Cell Treatments, and Synthetic Neural Scaffolds, Have Shown Promising Preclinical and Even Clinical Results with Potential Regenerative Treatment. <https://doi.org/10.31219/osf.io/m8cs9>.

Moataz Dowaidar. Photothermal and Photodynamic Photoactivation of Nanomaterials-Based Prodrugs Are Two Key Methods for NIR Light-Mediated Photoactivation. <https://doi.org/10.31219/osf.io/2bh3r>.

Moataz Dowaidar. Plant Viral Nanoparticles Can Be Used in Biological Systems for Loading and Transporting Cargo. <https://doi.org/10.31219/osf.io/txdka>.

Moataz Dowaidar. Polycomb Genes Role in Cancer Pathophysiology Is Offering Targets for Therapeutics Including Gene Therapy. <https://doi.org/10.31219/osf.io/sfvej>.

Moataz Dowaidar. Polydopamine May Be Easily Functionalized with a Range of Nanomaterials for Synergistic Cancer Therapy, in Addition to Its Exceptional Photothermal Effects. <https://doi.org/10.31219/osf.io/cq942>.

Moataz Dowaidar. Polydopamine Nanoparticles' Activity and Long-Term Stability Should Be Fully Studied for Gene Therapy Applications. <https://doi.org/10.31219/osf.io/x4nej>.

Moataz Dowaidar. Potential HIV Gene Therapy Strategies. <https://doi.org/10.31219/osf.io/e5hm2>.

Moataz Dowaidar. Potential Strategies for Cancer Gene Therapy. <https://doi.org/10.31219/osf.io/atcqz>.

Moataz Dowaidar. Potential Therapeutics for Primary Mitochondrial Disorders. <https://doi.org/10.31219/osf.io/6pz5k>.

Moataz Dowaidar. Potentials of Medicinal Nanostructured Diamond Particles and Coatings. <https://doi.org/10.31219/osf.io/h68xz>.

Moataz Dowaidar. Preclinical Investigations Revealed Possibilities for Salmonella Tumor Treatment. Bacteria Can Also Be Coupled to Nanomaterials Enabling Drug-Loading, Photocatalytic And/or Magnetic Properties, Using the Bacteria's Net Negative Charge. <https://doi.org/10.31219/osf.io/embqk>.

Moataz Dowaidar. Preclinical Studies and Clinical Trials Have Sparked Interest in Certain Biological Medications for Atherosclerotic Coronary Heart Disease. <https://doi.org/10.31219/osf.io/ts8mh>.

Moataz Dowaidar. Quantitative Groups Will Be Critical to the Success of Future Gene Therapy Programs. <https://doi.org/10.31219/osf.io/v97ht>.

Moataz Dowaidar. Quantum Dots Have the Potential to Be Used in Gene Therapy. <https://doi.org/10.31219/osf.io/bdeg6>.

Moataz Dowaidar. Research into P2X Purinergic Receptor Function in Tumor Growth Has Made Substantial Progress with Potential Gene Therapy Targeting. <https://doi.org/10.31219/osf.io/r34fs>.

Moataz Dowaidar. Research on Cell Sources for Brain Cell Replacement Methods Has Gained Major Importance. Cell and Gene Therapy Are Potentially Intriguing New Domains of Regenerative Medicine. <https://doi.org/10.31219/osf.io/g835b>.

Moataz Dowaidar. Researchers Would Be Able to Develop a Detailed Picture of Chromatin in Disease, Which Would Be Useful for Gene Therapy. <https://doi.org/10.31219/osf.io/m9z48>.

Moataz Dowaidar. RNA Sequencing and Microarray Analysis Are Helpful Techniques to Detect Obesity-Related lncRNAs. lncRNA Can Alter Cholesterol Metabolism and Can Be a Target for Gene Therapy. <https://doi.org/10.31219/osf.io/3fb6w>.

Moataz Dowaidar. RNA Therapies Hold Great Promise for Treating Cancer. High-Throughput Screening Techniques Have Facilitated the Development of RNA Treatments. <https://doi.org/10.31219/osf.io/9vxrb>.

Moataz Dowaidar. RNA-Based Gene Therapy for Manipulating the Neuroinflammatory Cascade Closely Linked to Neurodegeneration Can Help Reduce Disease Development. <https://doi.org/10.31219/osf.io/2hswv>.

Moataz Dowaidar. RNAi Treatment Has Been Shown to Successfully Modify Human-Related Target Gene Expression, Including Cancer. It Has the Capacity to Control Non-Standard Oncogenes, such as Oncogenic lncRNAs. <https://doi.org/10.31219/osf.io/bwqep>.

Moataz Dowaidar. RNAi-Based Gene Therapy Provides a Wide Variety of Applications. Safe, Biodegradable Nano Delivery Vectors Are Still Needed. <https://doi.org/10.31219/osf.io/s2zhn>.

Moataz Dowaidar. RNAs Hold a Lot of Potential When It Comes to Druggable Molecular Targets. <https://doi.org/10.31219/osf.io/2dtxg>.

Moataz Dowaidar. Sepsis-Associated Acute Kidney Damage Is a Disease That Affects the Patient's Quality of Life. It Should Be a Target for Gene Therapy. <https://doi.org/10.31219/osf.io/49k7q>.

- Moataz Dowaidar. Shadow Enhancers' Objective Seems to Be to Establish Robust Growth Patterns Independent of Genetic or Environmental Stress. <https://doi.org/10.31219/osf.io/qfinkp>.
- Moataz Dowaidar. Sickle Cell Disease Has Emerged as a Public Health Concern. Some Drugs May Conflict with Curative Therapies, yet They May Be Useful as a Bridge to HSCT and Gene Therapy. <https://doi.org/10.31219/osf.io/6kufh>.
- Moataz Dowaidar. Sickle Cell Disease Hematopoietic Stem Cell Gene Therapy with Globin Gene Addition Is Promising. <https://doi.org/10.31219/osf.io/j5fkb>.
- Moataz Dowaidar. Single-Gene Mutations in mtDNA-Associated Proteins Are Unlikely to Be the Main Cause of Sporadic Parkinson's Disease. Cumulative Genetic Variation in Numerous Genes May Be Important in Neurodegeneration and PD Risk. <https://doi.org/10.31219/osf.io/89qte>.
- Moataz Dowaidar. Small Nuclear Ribonucleoproteins (snRNPs) Based Gene Therapy. <https://doi.org/10.31219/osf.io/c43r9>.
- Moataz Dowaidar. Stimulator of Interferon Genes (STING)-Activating Nanoparticles Can Be Employed as a Tool for Controlled Immune Activation. <https://doi.org/10.31219/osf.io/2ez7a>.
- Moataz Dowaidar. Strategies for Treating Multiple Sclerosis with Gene Therapy. <https://doi.org/10.31219/osf.io/sync6>.
- Moataz Dowaidar. Studying the Pathologic Mechanisms of Osteoporosis and the Bone Microenvironment May Help Researchers Better Know the Etiology of Rheumatoid Arthritis, Periodontitis, and Multiple Myeloma, as Well as Other Inflammatory and Autoimmune Disorders. <https://doi.org/10.31219/osf.io/t3z6y>.
- Moataz Dowaidar. Suicide Gene Therapy May Be Effective in the Treatment of Malignant Glioma. <https://doi.org/10.31219/osf.io/vdkst>.
- Moataz Dowaidar. Synuclein Is a Protein That Is Expressed in Brain Tissue. The Specific Missense Mutation (SNCA) Found in a Family with Parkinson's Disease Is the Cause. Other Diseases Include Alzheimer's Disease and REM Sleep Behavior Disorder. <https://doi.org/10.31219/osf.io/bs8rc>.
- Moataz Dowaidar. Systems Biology Is a Method for Analyzing Massive Amounts of Multidimensional Data Generated by Omics Technologies. Cross-Validation of the Various Technological Platforms Is Critical. <https://doi.org/10.31219/osf.io/p8vkd>.
- Moataz Dowaidar. Targeted Chemical Nucleases Have a Wide Range of Untapped Applications in Biological Fields, Including Gene Therapy. <https://doi.org/10.31219/osf.io/6bexs>.
- Moataz Dowaidar. Targeting Mitochondria and Especially Taz Gene Mutation Induces CL May Give Novel Therapeutic Alternatives for Treating Barth Syndrome. <https://doi.org/10.31219/osf.io/unfpy>.
- Moataz Dowaidar. The Ability to Combine Multiple mRNA Antigens Targeting Multiple Pathogens Simultaneously, and the Robust Immune Responses Are Confirmed in Several Clinical Studies. <https://doi.org/10.31219/osf.io/6qksx>.
- Moataz Dowaidar. The Cardiometabolic-Based Chronic Disease Model Lays the Foundations for Accurate, Evidence-Based Preventive Targeting and Gene Therapy. <https://doi.org/10.31219/osf.io/up9z4>.
- Moataz Dowaidar. The Combination of Unique Biomolecules and Nanoparticles Has Shown Successful Gene Therapy Treatment Approaches for Non-Small Cell Lung Cancer Treatment. <https://doi.org/10.31219/osf.io/yeq5z>.
- Moataz Dowaidar. The Cubic Polyhedral Oligomeric Silsesquioxanes Based Hybrid Materials Have a Wide Variety of Applications, Including Drug Administration, Gene Therapy, Biological Imaging, and Bone Regeneration. <https://doi.org/10.31219/osf.io/9peq8>.
- Moataz Dowaidar. The Development of Tissue Replacement Therapies and Drug Discovery Was a Critical Milestone in Advancing Regenerative Medicine. <https://doi.org/10.31219/osf.io/w9bsm>.
- Moataz Dowaidar. The Epidemic of COVID-19 Prompted Widespread Use of mRNA Vaccinations. <https://doi.org/10.31219/osf.io/jqws5>.
- Moataz Dowaidar. The Gene Expression Profiling Gives an in-Depth Insight of Breast Cancer Heterogeneity, Better than a Single Protein or Gene Expression. It Is Time to Include It in the Daily Routine. <https://doi.org/10.31219/osf.io/xhyd7>.
- Moataz Dowaidar. The Most Useful and Commonly Available Acute Rejection Surveillance Strategies Are Routine Monitoring of Myocardial Function and Donor-Specific Anti-HLA Abs Monitoring. <https://doi.org/10.31219/osf.io/ebw68>.
- Moataz Dowaidar. The Nanomedicine System Has Successfully Inhibited Tumor Neovascularization Using Gene Silencing, Chemotherapy, Photothermal Therapy, and Other Therapies. <https://doi.org/10.31219/osf.io/rk2bf>.
- Moataz Dowaidar. The Protease MBTPS2 Is an Important Regulator of Several Cellular Processes, Especially in Health and Sickness. <https://doi.org/10.31219/osf.io/qyn6h>.
- Moataz Dowaidar. The Sigma 1 Receptor (S1R) Is a Potential Therapeutic Target for the Treatment of Huntington's Disease. <https://doi.org/10.31219/osf.io/mcefx>.
- Moataz Dowaidar. The Therapeutic Application of a Nucleic Acid Sequence to Patients' Diseased Organs Is Currently Available. <https://doi.org/10.31219/osf.io/pqsbf>.
- Moataz Dowaidar. The Treatment of Major Human Illnesses with Recombinant Adeno-Associated Virus (rAAV) Has Shown Tremendous Promises. <https://doi.org/10.31219/osf.io/uwa4e>.
- Moataz Dowaidar. The Use of a Network Medicine Approach Might Result in Innovative Strategies for Lowering Coronary Heart Disease and CV Risks. <https://doi.org/10.31219/osf.io/eakg8>.

- Moataz Dowaidar. The Vasoconstrictor Endothelin System Involvement in Chronic Kidney Diseases Pathogenesis Is Now the Most Often Employed Treatment Method. <https://doi.org/10.31219/osf.io/cnkqy>.
- Moataz Dowaidar. The VPS35-D620N Mutation Is Associated with Parkinson's Disease and Can Be a Target for Gene Therapy. <https://doi.org/10.31219/osf.io/83sxx>.
- Moataz Dowaidar. Therapeutics Including Gene Therapy for Osteoarthritis as a Concept. <https://doi.org/10.31219/osf.io/7zsqy>.
- Moataz Dowaidar. Thrombosis Pathways and Therapeutic Strategies. <https://doi.org/10.31219/osf.io/57vyz>.
- Moataz Dowaidar. Tissue Hypoxia Has Been Established as a Master Regulator for Alternative Splicing, with Substantial Clinical Consequences and Possibilities for Gene Therapy Targeting. <https://doi.org/10.31219/osf.io/5pbw4>.
- Moataz Dowaidar. To Rectify Alzheimer's Disease Etiology, Excessive Mitochondrial Division Might Be Stopped or Mitophagy Might Be Promoted. <https://doi.org/10.31219/osf.io/6kdxw>.
- Moataz Dowaidar. Transcriptomics Is a Rapidly Growing Field That Generates New Data That May Be Used on Its Own or in Combination with Existing Clinical Data for Development of New Therapeutics, Including Gene Therapy. <https://doi.org/10.31219/osf.io/kfr6a>.
- Moataz Dowaidar. Triple-Negative Breast Cancer, Which Lacks the Expression of Hormone Receptors and HER2, Has a Worse Prognosis. Massive Parallel Sequencing Is Capable of Reliably Breaking down the Intra-Tumor and Inter-Tumor Heterogeneity. <https://doi.org/10.31219/osf.io/pvk7u>.
- Moataz Dowaidar. Tumor Microenvironment Has Clinical Significance in Terms of Prognosis and Therapy Prediction. <https://doi.org/10.31219/osf.io/4dz8q>.
- Moataz Dowaidar. Tumor-Targeted Drug Delivery Systems for Anticancer Therapies Can Selectively Provide an Appropriate Cytotoxic Payload to Cancer Cells, Reducing the Side Effects of Chemo. <https://doi.org/10.31219/osf.io/683nj>.
- Moataz Dowaidar. Understanding Why the Same Gene Delivery Vector Behaves Differently in Different Cell Types Is Essential for Developing More Adaptable Transfection Systems. <https://doi.org/10.31219/osf.io/6q8af>.
- Moataz Dowaidar. Using AAV as a Gene Delivery Vector in the Neural System Is Effective in Several Animals, such as Nonhuman Primates. <https://doi.org/10.31219/osf.io/ut4fa>.
- Moataz Dowaidar. Using Pre-Existing Datasets to Combine Published Information with New Metrics Would Help Researchers Construct a Broader Picture of Chromatin in Disease. <https://doi.org/10.31219/osf.io/gsqv5>.
- Moataz Dowaidar. Virus-like Particles Are Good Nanocarriers for Liquid Biopsy Probes, Imaging Contrast Agents, and Anticancer Medications. <https://doi.org/10.31219/osf.io/xbtka>.
- Moataz Dowaidar. What Genomic Research Has Told Us about the Obesity and Its Possible Gene Therapy Targets. <https://doi.org/10.31219/osf.io/ym49s>.
- Moataz Dowaidar. ZEB1 Controls the Expression of ICAM1, Promoting Monocyte-Macrophage Adhesion and Hence the Formation of Atherosclerotic Lesions. <https://doi.org/10.31219/osf.io/kzjqg>.
- Nagahara AH, Mateling M, Kovacs I, Wang L, Eggert S, Rockenstein E, Koo EH, Masliah E, Tuszynski MH, 2013. Early BDNF treatment ameliorates cell loss in the entorhinal cortex of APP transgenic mice. *J. Neurosci* 33 (39), 15596–15602.
- Nagahara AH, Merrill DA, Coppola G, Tsukada S, Schroeder BE, Shaked GM, Wang L, Blesch A, Kim A, Conner JM, Rockenstein E, Chao MV, Koo EH, Geschwind D, Masliah E, Chiba AA, Tuszynski MH, 2009. Neuroprotective effects of brain-derived neurotrophic factor in rodent and primate models of Alzheimer's disease. *Nat. Med* 15 (3), 331–337.
- Nakaizumi T, Kawamoto K, Minoda R, Raphael Y, 2004. Adenovirus-mediated expression of brain-derived neurotrophic factor protects spiral ganglion neurons from ototoxic damage. *Audiol. Neuro. Otol* 9, 135–143. 10.1159/000077264.
- Nakajima H, Uchida K, Yayama T, Kobayashi S, Guerrero AR, Furukawa S, Baba H, 2010. Targeted retrograde gene delivery of brain-derived neurotrophic factor suppresses apoptosis of neurons and oligodendroglia after spinal cord injury in rats. *Spine* 35 (5), 497–504.
- Nam HJ, Gurda BL, McKenna R, Potter M, Byrne B, Salganik M, Muzyczka N, Agbandje-McKenna M, 2011. Structural studies of adeno-associated virus serotype 8 capsid transitions associated with endosomal trafficking. *J. Virol* 85, 11791–11799.
- Nathwani AC, Tuddenham EG, Rangarajan S, et al., 2011. Adenovirus-associated virus vector-mediated gene transfer in hemophilia B. *N. Engl. J. Med* 365, 2357–2365, 2011.
- Ortmann M, Zwitserlood P, Knief A, Baare J, Brinkheeter S, Am Zehnhoff-Dinnesen A, Dobel C, 2017. When hearing is tricky: speech processing strategies in prelingually deafened children and adolescents with cochlear implants having good and poor speech performance. *PloS One* 12 (1), e0168655 10.1371/journal.pone.0168655.
- Pan B, Askew C, Galvin A, Heman-Ackah S, Asai Y, Indzhykulian AA, Jodelka FM, Hastings ML, Lentz JJ, Vandenberghe LH, Holt JR, 2017. Géléc GS. Gene therapy restores auditory and vestibular function in a mouse model of Usher syndrome type 1c. *Nat. Biotechnol* 35 (3), 264–272.

- Panganiban LH, Barth JL, Darbelli L, Xing Y, Zhang J, Li H, Noble KV, Liu T, Brown LN, Schulte BA, Richard S, Lang H, 2018. Noise-induced dysregulation of quaking RNA binding proteins contributes to auditory nerve demyelination and hearing loss. *J. Neurosci* 2487–17.
- Pettingill LN, Richardson RT, Wise AK, O’Leary S, Shepherd RK, 2007. Neurotrophic factors and neural prostheses: potential clinical applications based upon findings in the auditory system. *IEEE Trans. Biomed. Eng* 54 (6), 1138–1148. 10.1109/TBME.2007.895375.
- Pfingst BE, Colesa DJ, Swiderski DL, Hughes AP, Strahl SB, Sinan M, Raphael Y, 2017. Neurotrophin gene therapy in deafened ears with cochlear implants: long-term effects on nerve survival and functional measures. *J Assoc Res Otolaryngol* 18 (6), 731–750. 10.1007/s10162-017-0633-9.
- Phillips HS, Hains JM, Laramée GR, Rosenthal A, Winslow JW, 1990. Widespread expression of BDNF but not NT3 by target areas of basal forebrain cholinergic neurons. *Science* 250 (4978), 290–294.
- Pirvola U, Ylikoski J, Palgi J, Lehtonen E, Arumäe U, Saarna M, 1992. Brain-derived neurotrophic factor and neurotrophin 3 mRNAs in the peripheral target fields of developing inner ear ganglia. *Proc. Natl. Acad. Sci. U S A* 89 (20), 9915–9919.
- Praetorius M, Brough DE, Hsu C, Plinkert PK, Pfannenstiel SC, Staecker H, 2008. Adenoviral vectors for improved gene delivery to the inner ear. *Hear. Res* 248, 31–38.
- Quintino L, Avallone M, Brannstrom E, Kavanagh P, Lockowandt M, Garcia Jareno P, Breger LS, Lindberg C, 2019. GDNF-mediated rescue of the nigrostriatal system depends on the degree of degeneration. *Gene Ther.* 26, 57–64. 10.1038/s41434-018-0049-0.
- Qun LX, Pirvola U, Saarna M, Ylikoski J, 1999. Neurotrophic factors in the auditory periphery. *Ann. N. Y. Acad. Sci* 884, 292–304.
- Rafii MS, Baumann TL, Bakay RA, Ostrove JM, Siffert J, Fleisher AS, Herzog CD, Barba D, Pay M, Salmon DP, Chu Y, Kordower JH, Bishop K, Keator D, Potkin S, Bartus RT, 2014. A phase I study of stereotactic gene delivery of AAV2-NGF for Alzheimer’s disease. *Alzheimers Dement* 10 (5), 571–581.
- Ramachandran PS, et al., 2017. Evaluation of dose and safety of AAV7m8 and AAV8BP2 in the non-human primate retina. *Hum. Gene Ther* 28, 154–167.
- Ramekers D, Versnel H, Grolman W, Klis SF, 2012. Neurotrophins and their role in the cochlea. *Hear. Res* 288 (1–2), 19–33. 10.1016/j.heares.2012.03.00.
- Ramekers D, Versnel H, Strahl SB, Klis SFL, Grolman W, 2015. Temporary neurotrophin treatment prevents deafness-induced auditory nerve degeneration and preserves function. *J. Neurosci* 35 (36), 12331–12345. 10.1523/JNEUROSCI.0096-15.2015.
- Rejali D, Lee VA, Abrashkin KA, Humayun N, Swiderski DL, Raphael Y, 2007. Cochlear implants and ex vivo BDNF gene therapy protect spiral ganglion neurons. *Hear. Res* 228, 180–187. 10.1016/j.heares.2007.02.010.
- Rio C, Dikkes P, Liberman MC, Corfas G, 2002. Glial fibrillary acidic protein expression and promoter activity in the inner ear of developing and adult mice. *J. Comp. Neurol* 442, 156–162.
- Rubel EW, Fritsch B, 2002. Auditory system development: primary auditory neurons and their targets. *Annu. Rev. Neurosci* 25, 51–101. 10.1146/annurev.neuro.25.112701.142849.
- Russell S, et al., Bennett J, Wellman JA, Chung DC, Yu ZF, Tillman A, 2017. Efficacy and safety of voretigene neparvovec (AAV2- hrPE65v2) in patients with RPE65-mediated inherited retinal dystrophy: a randomised, controlled, open-label, phase 3 trial. *Lancet* 390, 849–860. 10.1016/S0140-6736(17)31868-8. Epub 2017 Jul 14.
- Sampaio TB, Savall AS, Gutierrez MEZ, Pinton S, 2017. Neurotrophic factors in Alzheimer’s and Parkinson’s diseases: implications for pathogenesis and therapy. *Neural .Regen. Res* 12 (4), 549–557.
- Schaette R, McAlpine D, 2011. Tinnitus with a normal audiogram: physiological evidence for hidden hearing loss and computational model. *J. Neurosci. : the official journal of the Society for Neuroscience* 31 (38), 13452–13457. 10.1523/JNEUROSCI.2156-11.2011.
- Schechterson LC, Bothwell M, 1994. Neurotrophin and neurotrophin receptor mRNA expression in developing inner ear. *Hear. Res* 73 (1), 92–100.
- Scheperle RA, 2017. Suprathreshold compound action potential amplitude as a measure of auditory function in cochlear implant users. *J. Otolaryngol* 12 (1), 18–28. 10.1016/j.joto.2017.01.001.
- Schwartz-Leyzac KC, Pfingst BE, 2018. Assessing the relationship between the electrically evoked compound action potential and speech recognition abilities in bilateral cochlear implant recipients. *Ear Hear.* 39, 344–358. 10.1097/AUD.0000000000000490.

Sereyenko et al., 2013.

Settin, Ahmad A., Abdullah Algasham, Moataz Dowaidar, and Hisham Ismail. 2009. Methylene Tetrahydrofolate Reductase and Angiotensin Converting Enzyme Gene Polymorphisms Related to Overweight/obesity among Saudi Subjects from Qassim Region. *Disease Markers* 27 (2): 97–102. <https://doi.org/10.3233/DMA-2009-0660>.

Settin, Ahmad A., Abdullah Alghasham, Ahmad Ali, Moataz Dowaidar, and Hisham Ismail. 2012. Frequency of Thrombophilic Genetic Polymorphisms among Saudi Subjects Compared with Other Populations. *Hematology* 17 (3): 176–82. <https://doi.org/10.1179/102453312X13376952196575>.

Settin, Ahmad, Abdullah Algasham, Moataz Dowaidar, and Hisham Ismail. 2011. Methylene Tetrahydrofolate Reductase (MTHFR) and Angiotensinogen Converting Enzyme (ACE) Gene Polymorphisms Related to Overweight and Obesity among Saudi Patients in Al Qassim. *International Journal of Health Sciences* 5 (2 Suppl 1): 24–25. <https://www.ncbi.nlm.nih.gov/pubmed/23284565>.

Settin, Ahmad, Hala Almarsafawy, Ahmad Alhussieny, and Moataz Dowaidar. 2008a. Dysmorphic Features, Consanguinity and Cytogenetic Pattern of Congenital Heart Diseases: A Pilot Study from Mansoura Locality, Egypt. *International Journal of Health Sciences* 2 (2): 101–11. <https://www.ncbi.nlm.nih.gov/pubmed/21475491>.

Settin, Ahmad, Ibrahim S. Abu-Saif, Rizk El-Baz, Moataz Dowaidar, Rabab Abu-Al Kasim, and Shaimaa Shabana. 2007a. Diagnosis of Sex Chromosome Disorders and Prenatal Diagnosis of Down Syndrome Using Interphase Fluorescent In-Situ Hybridization Technique. *International Journal of Health Sciences* 1 (2): 203–9. <https://www.ncbi.nlm.nih.gov/pubmed/21475429>.

Settin, Ahmad, Moataz Dowaidar, Rizk El-Baz, Ayman Abd-Al-Samad, Ibrahim El-Sayed, and Mahmoud Nasr. 2008. Frequency of Factor V Leiden Mutation in Egyptian Cases with Myocardial Infarction. *Hematology* 13 (3): 170–74. <https://doi.org/10.1179/102453308X316158>.

Seyyedi M, Viana LM, Nadol JB Jr., 2014. Within-subject comparison of word recognition and spiral ganglion cell count in bilateral cochlear implant recipients. *Otol. Neurotol* 35 (8), 1446–1450. 10.1097/MAO.0000000000000443.

Shepherd RK, Coco A, Epp SB, 2008. Neurotrophins and electrical stimulation for protection and repair of spiral ganglion neurons following sensorineural hearing loss. *Hear. Res* 242, 100–109. 10.1016/j.heares.2007.12.005.

Shepherd RK, Coco A, Epp SB, Crook JM, 2005. Chronic depolarization enhances the trophic effects of brain-derived neurotrophic factor in rescuing auditory neurons following a sensorineural hearing loss. *J. Comp. Neurol* 486, 145–158.

Shibata SB, Cortez SR, Beyer LA, Wiler JA, Di Polo A, Pflugst BE, Raphael Y, 2010. Transgenic BDNF induces nerve fiber regrowth into the auditory epithelium in deaf cochleae. *Exp. Neurol* 223 (2), 464–472.

Shibata SB, Di Pasquale G, Cortez SR, Chiorini JA, Raphael Y, 2009. Gene transfer using bovine adeno-associated virus in the Guinea pig cochlea. *Gene Ther.* 16 (8), 990–997.

Shibata SB, Osumi Y, Yagi M, Kanda S, Kawamoto K, Kuriyama H, Nishiyama T, Yamashita T, 2007. Administration of amitriptyline attenuates noise-induced hearing loss via glial cell line-derived neurotrophic factor (GDNF) induction. *Brain Res.* 1144, 74–81. 10.1016/j.brainres.2007.01.090.

Shoji F, Yamasoba T, Magal E, Dolan DF, Altschuler RA, Miller JM, 2000. Glial cell line-derived neurotrophic factor has a dose dependent influence on noise-induced hearing loss in the Guinea pig cochlea. *Hear. Res* 142 (1–2), 41–55. 10.1016/S0378-5955(00)00007-1.

Simmons DA, Longo FM, Massa SM, 2017. Neurotrophin receptor signaling as a therapeutic target for Huntington’s disease. *CNS Neurol. Disord. - Drug Targets* 16 (3), 291–302.

Simonelli F, Maguire AM, Testa F, et al., 2010. Gene therapy for Leber’s congenital amaurosis is safe and effective through 1.5 years after vector administration. *Mol. Ther* 18, 643–650.

Sly DJ, Campbell L, Uschakov A, Saief ST, Lam M, O’Leary SJ, 2016. Applying neurotrophins to the round window rescues auditory function and reduces inner hair cell synaptopathy after noise-induced hearing loss. *Otol. Neurotol* 37 (9), 1223–1230. 10.1097/MAO.0000000000001191.

Spoendlin, 1984. Factors inducing retrograde degeneration of the cochlear nerve. *Ann. Otol. Rhinol. Laryngol. Suppl* 112, 76–82.

Staecker H, Jolly C, Garnham C, 2010. Cochlear implantation: an opportunity for drug development. *Drug Discov. Today* 15, 314–321. 10.1016/j.drudis.2010.02.005.

Staecker H, Kopke R, Malgrange B, Lefebvre P, Van de Water TR, 1996. NT-3 and/or BDNF therapy prevents loss of auditory neurons following loss of hair cells. *Neuroreport* 22 (7), 889–894, 4.

Staecker H, Schlecker C, Kraft S, Praetorius M, Hsu C, Brough DE, 2014. Optimizing atoh1-induced vestibular hair cell regeneration. *Laryngoscope* 124, S1–S12.

- Stankovic K, Rio C, Xia A, Sugawara M, Adams JC, Liberman MC, Corfas G, 2004. Survival of adult spiral ganglion neurons requires erbB receptor signaling in the inner ear. *J. Neurosci* 24 (40), 8651–8661.
- Sugawara M, Murtie JC, Stankovic KM, Liberman MC, Corfas G, 2007. Dynamic patterns of neurotrophin 3 expression in the postnatal mouse inner ear. *J. Comp. Neurol* 501 (1), 30–37.
- Suzuki J, Corfas G, Liberman MC, 2016. Round-window delivery of neurotrophin 3 regenerates cochlear synapses after acoustic overexposure. *Sci. Rep* 6, 24907 10.1038/srep24907.
- Suzuki J, Hashimoto K, Xiao R, Vandenberghe LH, Liberman MC, 2017. Cochlear gene therapy with ancestral AAV in adult mice: complete transduction of inner hair cells without cochlear dysfunction. *Sci. Rep* 7, 45524 10.1038/srep45524. Erratum. *Sci Rep*. 2017 May 22;7:46827.
- Svirsky MA, Robbins AM, Kirk KI, Pisoni DB, Miyamoto RT, 2000. Language development in profoundly deaf children with cochlear implants. *Psychol. Sci* 11, 153–158.
- Tagoe T, Barker M, Jones A, Allcock N, Hamann M, 2014. Auditory nerve perinodal dysmyelination in noise-induced hearing loss. *J. Neurosci* 34 (7), 2684–2688.
- Tao Y, Huang M, Shu Y, Ruprecht A, Wang H, Tang Y, Vandenberghe LH, Wang Q, Gao G, Kong WJ, Chen ZY, 2018. Delivery of adeno-associated virus vectors in adult mammalian inner-ear cell subtypes without auditory dysfunction. *Hum. Gene Ther* 29 (4), 492–506. 10.1089/hum.2017.120.
- Thanos C, Emerich D, 2005. Delivery of neurotrophic factors and therapeutic proteins for retinal diseases. *Expert Opin. Biol. Ther* 5, 1443–1452. Review.
- Tuszynski MH, Thal L, Pay M, Salmon DP, Bakay R, Patel P, Blesch A, Vahlsing HL, Ho G, Tong G, Potkin SG, Fallon J, Hansen L, Mufson EJ, Kordower JH, Gall C, Conner J, 2005. A phase 1 clinical trial of nerve growth factor gene therapy for Alzheimer disease. *Nat. Med* 11 (5), 551–555.
- Tuszynski MH, Yang JH, Barba D, U HS., Bakay RA, Pay MM, Masliah E, Conner JM, Kobalka P, Roy S, Nagahara AH, 2015. Nerve growth factor gene therapy: activation of responses and Alzheimer disease, neuronal responses in Alzheimer disease. *JAMA Neurol* 72, 1139–1147. 10.1001/jamaneurol.2015.1807.
- Uchida K, Nakajima H, Hirai T, Yayama T, Chen K, Guerrero AR, Johnson WE, Baba H, 2012. The retrograde delivery of adenovirus vector carrying the gene for brain-derived neurotrophic factor protects neurons and oligodendrocytes from apoptosis in the chronically compressed spinal cord of *twy/twy* mice. *Spine* 37 (26), 2125–2135. 10.1097/BRS.0b013e3182600ef7.
- Vandenberghe LH, et al., 2011. Dosage thresholds for AAV2 and AAV8 photoreceptor gene therapy in monkey. *Sci. Transl. Med* 3, 88ra54.
- Venit, Tomas, Moataz Dowaidar, Maxime Gestein, Syed Raza Mahmood, Ülo Langel, and Piergiorgio Percipalle. 2020. Transcriptional Profiling Reveals Ribosome Biogenesis, Microtubule Dynamics and Expression of Specific lncRNAs to Be Part of a Common Response to Cell-Penetrating Peptides. *Biomolecules* 10 (11): 1567. <https://doi.org/10.3390/biom10111567>.
- Ventriglia M, Zanardini R, Bonomini C, Zanetti O, Volpe D, Pasqualetti P, Gennarelli M, Bocchio-Chiavetto L, 2013. Serum brain-derived neurotrophic factor levels in different neurological diseases. *BioMed Res. Int* 901082.
- Völter C, Götze L, Dazert S, Falkenstein M, Thomas JP, 2018. Can cochlear implantation improve neurocognition in the aging population? *Clin. Interv. Aging* 13, 701–712. 10.2147/CIA.S160517.
- Wan G, Gómez-Casati ME, Gigliello AR, Liberman MC, Corfas G, 2014. Neurotrophin-3 regulates ribbon synapse density in the cochlea and induces synapse regeneration after acoustic trauma. *Elife* 20, 3 10.7554/eLife.03564.
- Wang Q, Green SH, 2011. Functional role of neurotrophin-3 in synapse regeneration by spiral ganglion neurons on inner hair cells after excitotoxic trauma in vitro. *J. Neurosci* 31 (21), 7938–7949. 10.1523/JNEUROSCI.1434-10.2011.
- Watakabe A, Ohtsuka M, Kinoshita M, Takaji M, Isa K, Mizukami H, Ozawa K, Isa T, Yamamori T, 2015. Comparative analyses of adeno-associated viral vector serotypes 1, 2, 5, 8 and 9 in marmoset, mouse and macaque cerebral cortex. *Neurosci. Res* 93, 144–157. 10.1016/j.neures.2014.09.002.
- Weishaupt N, Blesch A, Fouad K, 2012. BDNF: the career of a multifaceted neurotrophin in spinal cord injury. *Exp. Neurol* 238 (2), 254–264. 10.1016/j.expneurol.2012.09.001.
- WHO, 2019. World Health Organization. <https://www.who.int/news-room/fact-sheets/detail/deafness-and-hearing-loss>. [Google Scholar]

- Wise AK, Hume CR, Flynn BO, Jeelall YS, Suhr CL, Sgro BE, O'Leary SJ, Shepherd RK, Richardson RT, 2010. Effects of localized neurotrophin gene expression on spiral ganglion neuron resprouting in the deafened cochlea. *Mol. Ther* 18, 1111–1122. 10.1038/mt.2010.28.
- Wise AK, Richardson R, Hardman J, Clark G, O'Leary S, 2005. Resprouting and survival of Guinea pig cochlear neurons in response to the administration of the neurotrophins brain-derived neurotrophic factor and neurotrophin-3. *J. Comp. Neurol* 487, 147–165. 10.1002/cne.20563.
- Wise AK, Tu T, Atkinson PJ., Flynn BO, Sgro BE, Hume C, O'Leary SJ, Shepherd RK, Richardson RT, 2011. The effect of deafness duration on neurotrophin gene therapy for spiral ganglion neuron protection. *Hear. Res* 278, 69–76. 10.1016/j.heares.2011.
- Won JH, Drennan WR, Kang RS, Rubinstein JT, 2010. Psychoacoustic abilities associated with music perception in cochlear implant users. *Ear Hear.* 31, 796–805. 10.1097/AUD.0b013e3181e8b7bd.
- Wu J, Liu B, Fan J, Zhu Q, Wu J, 2011. Study of protective effect on rat cochlear spiral ganglion after blast exposure by adenovirus- mediated human β -nerve growth factor gene. *Am. J. Otolaryngol* 32, 8–12. 10.1016/j.amjoto.2009.08.012.
- Xia L, Yin S, Wang J, 2012. Inner ear gene transfection in neonatal mice using adeno-associated viral vector: a comparison of two approaches. *PloS One* 7, e43218 10.1371/journal.pone.0043218.
- Xiong W, Wu DM, Xue Y, Wang SK, Chungc MJ, Ji X, Rana P, Zhao SR, Maia S, Cepko CL, 2019. AAV cis-regulatory sequences are correlated with ocular toxicity. *Proc. Natl. Acad. Sci. Unit. States Am* 116 (12), 5785–5794. 10.1073/pnas.1821000116.
- Yagi M, Kanzaki S, Kawamoto K, Shin B, Shah PP, Magal E, Sheng J, Raphael Y, 2000. Spiral ganglion neurons are protected from degeneration by GDNF gene therapy. *J Assoc Res Otolaryngol* 1, 315–325.
- Yang T, Kersigo J, Jahan I, Pan N, Fritzs B, 2011. The molecular basis of making spiral ganglion neurons and connecting them to hair cells of the organ of Corti. *Hear. Res* 278, 21–33. 10.1016/j.heares.2011.03.002.
- Ylikoski J, Pirvola U, Moshnyakov M, Palgi J, Arumae U, Saarma M, 1993. Expression patterns of neurotrophin and their receptor mRNAs in the rat inner ear. *Hear. Res* 65, 69–78.
- Ylikoski J, Pirvola U, Virkkala J, Suvanto P, Liang XQ, Magal E, Altschuler R, Miller JM, Saarma M, 1998. Guinea pig auditory neurons are protected by glial cell line-derived growth factor from degeneration after noise trauma. *Hear. Res* 124, 17–26.
- Yu-Wai-Man P, Votruba M, Moore AT, Chinnery PF, 2014. Treatment strategies for inherited optic neuropathies: past, present and future. *Eye* 28, 521–537. 10.1038/eye.2014.37.
- Zeng F-G, Rebscher S, Harrison W, Sun X, Fen H, 2008. Cochlear implants: system design, integration, and evaluation. *IEEE Rev Biomed Eng* 1, 115–142. 10.1109/RBME.2008.2008250.
- Zhao Y, Li Y, Zheng Z, Li J, Nie X, Jin X, Zheng J, Zhang J, Chen M, Hao J, Yang Y, Liu W, Liu H, Ni X, 2019. Health-related quality of life in Mandarin-speaking children with cochlear implants. *Ear Hear.* 40 (3), 605–614. 10.1097/AUD.0000000000000633.
- Ziemlińska E, Kügler S, Schachner M, Wewiór I, Czarkowska-Bauch J, Skup M, 2014. Overexpression of BDNF increases excitability of the lumbar spinal network and leads to robust early locomotor recovery in completely spinalized rats. *PloS One* 9 (2), e88833.