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New approaches to the treatment of orphan genetic disorders:
Mitigating molecular pathologies using chemicals

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ABSTRACT

With the advance and popularization of molecular techniques, the identification of genetic mutations that cause diseases has increased dramatically. Thus, the number of laboratories available to investigate a given disorder and the number of subsequent diagnosis have increased over time. Although it is necessary to identify mutations and provide diagnosis, it is also critical to develop specific therapeutic approaches based on this information. This review aims to highlight recent advances in mutation-targeted therapies with chemicals that mitigate mutational pathology at the molecular level, for disorders that, for the most part, have no effective treatment. Currently, there are several strategies being used to correct different types of mutations, including the following: the identification and characterization of translational readthrough compounds; antisense oligonucleotide-mediated splicing redirection; mismatch repair; and exon skipping. These therapies and other approaches are reviewed in this paper.

Key words: antisense oligonucleotide-mediated splicing redirection, mismatch repair, mutation-targeted therapies, translational readthrough.

INTRODUCTION

Many human diseases have an identified genetic basis. Whereas some of these diseases may be attributed to mutations at multiple loci, a significant number of diseases are believed to be caused by mutations in single genes – monogenic diseases. These monogenic disorders can be subdivided into autosomal dominant, autosomal recessive and X-linked diseases. Although each of these conditions is relatively rare, the many different types of monogenic diseases together affect a substantial portion of the population. The World Health Organization (WHO 2014) estimated the
global prevalence of all single gene diseases to be approximately 10 in every 1000 births. Some of these diseases can be very severe and even fatal (Wong and Chiu 2010).

Genetic mutations can be classified into four major groups: missense, nonsense, splicing and frameshift. Missense mutations are typically single nucleotide changes that either alter the amino acid in translated proteins (nonsynonymous variations) or do not alter the amino acid (synonymous or ‘silent’ variations). Nonsense mutations are point mutations in a sequence that create a UAA, UAG, or UGA codon in the coding region of the mRNA, resulting in a premature translation termination and, usually, a nonfunctional or rapidly degraded protein. Splicing mutations result in disruption of critical sequences for splicing, and abolishment of the usual splice sites, or creation of aberrant or cryptic splice sites, which in turn resulting in aberrant proteins. Frameshift mutations most commonly are caused by deletion or insertion of a number of nucleotides that alter the reading frame for any subsequent downstream codons (Hu and Gatti 2008, Orro et al. 2008, Mitui et al. 2009).

The existence of common groups of mutations prompted the hypothesis of common group-specific molecular pathogenesis, e.g., that therapeutic strategies developed against a common mutation group will be effective against similar mutations, regardless of the gene (Hu and Gatti 2008). In this review, we highlight mutation-targeted therapies with chemicals that mitigate mutational pathology at the molecular level (Fig. 1), mainly for recessive disorders for which there is no effective treatment (Table I).

Figure 1 - Mutation-targeted therapies reviewed in this paper.
Mitigating Molecular Pathology Using Chemicals

Mutations that introduce premature stop codons (PTCs) into the coding sequences of genes have been implicated in numerous inherited diseases and several cancers, and at least 2,400 different genetic disorders have at least one causative nonsense allele (Mort et al. 2008, OMIM 2015).

In general, it has been estimated that nonsense mutations account for ~11% of all variations that cause inherited disorders and as many as 20% of the protein-coding region located, single base-pair mutations that cause these diseases (Ryan 2014, HGMD 2015). PTCs can originate from nonsense mutations, frameshift mutations, or from aberrant splicing that generates mRNA isoforms that lead to impaired translation.

### Examples of Mutation-Targeted Therapies

<table>
<thead>
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<th>Target*</th>
<th>Diseases Tested</th>
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<td>Diop et al. 2007, Nudelman et al. 2009, Goldmann et al. 2010</td>
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</tr>
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*The therapies are not exclusive to the targets indicated in this table; they could be used for other targets. (–) Not tested.

### AMINOGLYCOSIDES AND OTHERS DRUGS THAT PROMOTE STOP CODON READTHROUGH

In general, it has been estimated that nonsense mutations account for ~11% of all variations that cause inherited disorders and as many as 20% of the protein-coding region located, single base-pair mutations that cause these diseases (Ryan 2014, HGMD 2015). PTCs can originate from nonsense mutations, frameshift mutations, or from aberrant splicing that generates mRNA isoforms that lead...
to the production of nonfunctional, truncated or deleterious proteins (Linde and Kerem 2008, Kandasamy et al. 2011). Typically, these mutations result in the activation of nonsense mediated RNA decay (NMD), whereby PTC-containing mRNAs are targeted for rapid degradation. This mechanism blocks the production of truncated proteins, which leads to dominant-negative effects on cell function (Silva and Romão 2009, Almeida et al. 2012). Strategies for inducing stop codon readthrough offer the possibility of reducing the pathology caused by this type of mutation. There are at least two different scenarios in which PTC therapies may be maximally effective: (i) diseases in which a much lower than normal level of the deficient protein activity may be therapeutic (e.g., hemophilia and cystic fibrosis - CF); and (ii) diseases in which the therapeutic protein has a very long half-life and thus can accumulate to significant levels via PTC suppression (e.g., Duchene muscular dystrophy - DMD) (Hainrichson et al. 2008, Kandasamy et al. 2011).

Aminoglycosides such as gentamicin, amikacin and tobramycin have important clinical applications in the treatment of serious Gram-negative bacterial infections, and in the experimental treatment of recessive diseases with nonsense mutations (Jana and Deb 2006, Hainrichson et al. 2008, Vecsler et al. 2011, Peltz et al. 2013). These molecules bind to the decoding site of the 16S or 18S ribosomal RNA in prokaryotes and eukaryotes, respectively, inducing a local conformational change (Hu and Gatti 2008, Halvey et al. 2012). In prokaryotes, the binding of aminoglycoside is highly specific and is mediated through adenine 1408 in the 16S ribosome, thus leading to the inhibition of protein synthesis. However, in eukaryotes, the corresponding nucleotide is a guanine and the binding is less efficient, resulting in translational readthrough by the insertion of an amino acid at the stop codon (Keeling and Bedwell 2002, Sánchez-Alcudia et al. 2012).

Aminoglycoside antibiotics were the first small-molecule drugs that rendered promising results, and this approach has been validated by numerous in vitro and in vivo experiments in different disorders, including CF, DMD, Hurler’s syndrome (MPS I), and diabetes, among others (Diop et al. 2007, Nudelman et al. 2009, Goldmann et al. 2010). With direct clinical relevance, some trials of aminoglycoside therapy have been carried out in humans with PTC mutations. The most promising resulted from topically administered gentamicin in CF patients. After treatment, nasal potential difference measurements improved considerably in individuals with PTC mutations. Furthermore, full-length CFTR protein was detected in the nasal epithelial cells of two treated individuals (Wilschanski et al. 2003, Clancy et al. 2006, Sermet-Gaudelus et al. 2007). In patients with nonsense mutation-mediated DMD, there was evidence that intravenous gentamicin administration suppressed nonsense mutations, as determined by the analysis of full-length dystrophin in muscle biopsies (Politano et al. 2003). These experiments demonstrated the ability of selected aminoglycoside structures to induce mammalian ribosomes to read through disease-causing PTCs (Fig. 2) and partially restore full-size functional proteins (Burke and Mogg 1985, Kandasamy et al. 2011, Sánchez-Alcudia et al. 2012).

However, readthrough efficiency depends on the PTC (UGA > UAG > UAA); on the nucleotide context of the mutation, particularly the nucleotide immediately after the stop codon (C > U > A ≥ G); and on other factors involved in readthrough regulation (Bidou et al. 2004, Kimura et al. 2005, Diop et al. 2007). In addition, the efficiency of aminoglycoside therapy depends on the molecular mechanism by which the nonsense mutation cause the defects and on the level of active protein required to reduce clinical symptoms. Moreover, results are impaired by the need for regular intravenous administration and the toxic side effects (nephrotoxicity and ototoxicity) of long-term treatment (Finkel 2010). With the aim of identifying alternative compounds that do not elicit toxic side effects, several studies have been
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Figure 2 - Schematic diagram of gene regulation and readthrough strategy. The relative positions of caps, polyadenylation sequences, and initiation and termination codons within mRNAs are shown. (a) In wild type cells, normal mRNAs encode complete proteins. (b) In mutant cells, the mRNAs contain a premature stop codon, and truncated proteins are translated in the absence of aminoglycoside or the transcript is degraded by the NMD pathway. The presence of aminoglycoside allows the incorporation of a random amino acid at the terminal stop codon of the mutant mRNA. Full-length proteins result from the aminoglycoside-induced readthrough.

results for DMD, other dystrophinopathies, CF, MPS I, and Carnitine Palmitoyltransferase 1A Deficiency (CPT1A), with the potential of treating a wider range of genetic disorders (Finkel 2010, Sermet-Gaudelus et al. 2010, Rowe et al. 2012, Peltz et al. 2013, Ryan 2014).

A set of clinical studies have been conducted with ataluren in CF and DMD. A phase IIb international, randomized, double blind, placebo-controlled study evaluated the effects of ataluren therapy on ambulatory ability in 174 patients (≥5 years of age) with DMD for 48 weeks (Finkel et al. 2010, 2013, McDonald et al. 2013a, b). Patients
were stratified based on their age (<9 versus ≥9 years), use of corticosteroids (yes versus no), and baseline 6-min walk distance (6MWD) (<350 or ≥350 m) and were randomized 1:1:1 to placebo, ataluren low dose, or ataluren high dose with all regimens given three times per day. This study showed that ataluren 10, 10, 20 mg/kg was more effective than placebo, increasing the mean 6MWD by 31.3 m over the 48-week period and representing a 48% reduction in the risk of the 6MWD worsening by 10%. Timed function tests of muscle function (walking/running 10 m, up and down stairs) also revealed positive trends for ataluren 10, 10, 20 mg/kg, as evidenced by less decline over the 48 weeks. Ataluren showed activity and safety in this short-term study, supporting the further evaluation of ataluren 10, 10, 20 mg/kg and 20, 20, 40 mg/kg in similar future studies (Finkel et al. 2013, McDonald et al. 2013a, b).

In CF patients ≥ 6 years of age, a phase III 48-week, double-blind study was performed to assess the safety and efficacy of ataluren (Clancy et al. 2006, Rowe et al. 2012, Kerem et al. 2014). Patients were stratified by age, chronic inhaled antibiotic use and percent-predicted forced expiratory volume in 1 s (FEV1). At Week 48, the difference in the mean relative change from baseline in percent-predicted FEV1 between ataluren and placebo was 3.0%, and the mean pulmonary exacerbation rate was 23% lower for ataluren (Peltz et al. 2013, Kerem et al. 2014). These results were more important in patients who did not chronically take inhaled antibiotics. In this study, several inhaled antibiotics were used chronically by patients, including colistin, aztreonam and tobramycin. However, analysis of the effects of these different inhaled antibiotics on percent-predicted FEV1 and pulmonary exacerbation rates indicated that tobramycin antagonized the effect of ataluren, a finding which was confirmed by in vitro assays. The safety profiles were similar for ataluren and placebo (Rowe et al. 2012, Peltz et al. 2013, Kerem et al. 2014).

PTC124/ataluren (Translarna™ – PTC Therapeutics, Inc.) is a small molecule, orally available compound (Finkel et al. 2013, Ryan 2014), which is the first drug developed specifically to treat diseases caused by nonsense mutations. Notably, a conditional first approval for the use in European of this drug was achieved for DMD in 2014 (Ryan 2014, EMA 2015).

### USE OF THERAPEUTIC OLIGONUCLEOTIDES IN SPlicing MUTATIONS

RNA mis-splicing diseases account for up to 15% of all inherited diseases, ranging from neurological to myogenic to metabolic disorders. With the great increase in the performance of genomic sequencing for individual patients, the number of known mutations that affect splicing has risen to 50–60% of all disease-causing mutations (Wang and Cooper 2007, Baralle et al. 2009, Hammond and Wood 2011). Thus, correction or redirection of pre-mRNA splicing in a mutation-specific context represents a potential gene therapy modality with applicability to many inherited disorders.

Splicing mutations can be grouped into a minimum of five types. Type I is the classical splicing mutation that results in the deletion of an entire exon. Types II, III, and IV are nonclassical splicing mutations that result in pseudoexon inclusion or partial exon deletion. Type V involves the branch point; although such mutations exist, they have not been targeted for therapy (Hu and Gatti 2008). Exon inclusion, exon exclusion and exon skipping to restore the open reading frame and cryptic splicing are techniques used to correct frameshift mutations (Spitali and Aartsma-Rus 2012).

Antisense oligonucleotide (AON) therapies have been used to correct pre-mRNA splicing in many disease models. Modified oligonucleotides can alter the pathogenic splice signals that are activated by mutations, resulting in close to normal levels of mRNAs that encode functional protein; however, the therapeutic principles for each AON varies considerably (Fig. 3 - Wilton and Fletcher 2011).
The chemicals that are used to work in animal models include peptide nucleic acids (PNAs), alternating locked nucleic acids (LNAs) and deoxynucleotide oligonucleotides, fully modified (non-gapmer) 2'-substituted oligonucleotides and phosphorodiamidate morpholino (PMO)-based antisense oligonucleotides (aoNs) therapies are used to correct pre-mRNA splicing.

**Figure 3 - Antisense oligonucleotides (AONs) therapies are used to correct pre-mRNA splicing.**

(a) Exon exclusion: Large-scale deletions can result in out-of-frame mRNA transcripts. AONs directed toward an exon splice enhancer element within the mutated exon result in exclusion of the exon and, thus, an alternative in-frame transcript. (b) Exon inclusion: A mRNA transcript can lose one exon owing to negative splicing regulatory elements within and surrounding this exon and produce a less functional protein. Treatment with AON directed toward an intron splice silencer (ISS) can increase the inclusion of the exon from 20% to approximately 95%, as seen in the SMN2 gene (Hua et al. 2010). (c) Cryptic splice sites: This can result in inclusion of introns or partial deletions of exons from mRNA. The hypothetical intronic mutation illustrated creates an intronic 5' splice site (ss) and activates a cryptic 3' ss within intron 2. AONs targeting the mutated intron 5' ss blocks recognition by the splicing machinery and generates transcripts without the aberrant intron element. Similarly, a mutation within exon 2 creates a cryptic 5' ss preferentially utilized over the natural splice site. AONs directed to the mutant cryptic site cause steric blockage of the splicing machinery and redirect splicing to the natural 5' ss. (d) Pseudoexon: A C–A mutation within intron 1 strengthened the 50ss of a pseudoexon (dashed box), which maintains a strong 3' ss. AONs directed to the pseudoexon 5' ss and inclusion of C–A mutation restored the natural transcript (Adapted from Hammond and Wood 2011).
oligomers (Kole et al. 2012). By sterically blocking mutations and regulatory sequences within the pre-mRNA transcript, AONs have been used to induce the skipping or inclusion of exons, to block pseudoxons from recognition and to influence the alternative splicing of pathogenic isoforms (Hammond and Wood 2011).

Antisense oligonucleotides have been used to successfully modulate RNA splicing in CF (CFTR gene) (Friedman et al. 1999), β-thalassemia (β-globin gene) (Suwanmanee et al. 2002), Hutchinson–Gilford progeria syndrome (LMNA gene) (Scaffidi and Misteli 2005) and DMD (Dystrophin gene) (van Deutekom et al. 2007, Kinali et al. 2009, Cirak et al. 2011). These compounds have also been used to modify alternative splicing in the SMN2 (spinal muscular atrophy) (Hua et al. 2010, Nlend Nlend et al. 2010), Bcl-x (Taylor et al. 1999), and C-myc genes (Giles et al. 1999), suggesting that such antisense oligonucleotides are capable of crossing the blood–brain barrier in amounts sufficient to restore neural activity.

The data from the first clinical trial in DMD patients were highly encouraging, and technical advances in the development of methods for both directed exon exclusion and directed exon inclusion suggest that such antisense oligonucleotides are capable of crossing the blood–brain barrier in amounts sufficient to restore neural activity.

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Gene editing using ssODNs can alter single nucleotides and induce stable alterations at the genomic level. Several studies have tested technique optimizations. ssODNs are oligonucleotides that are < 200 bp in length and are designed to anneal to the lagging strand to generate a 100-fold greater ‘editing’ efficiency than those that anneal to the leading strand. The majority of editing events (∼70%) occur by the incorporation of the ssODN within the lagging strand during replication (Falgowski et al. 2011). Bertoni et al. (2009) showed that ssODNs that contain a methyl-CpG modification and are capable of binding to the methyl-CpG binding domain protein 4 (MBD4) are able to induce > 10-fold higher levels of gene correction than ssODNs lacking these specific modifications.

RDOs are a class of oligonucleotides used for gene targeting with a correction efficiency of approximately 50%; however, the reproducibility of these studies has been limited in mammalian systems (Sargent et al. 2011). TFOs are ssODNs that are typically 10–40 nt in length and bind to specific regions in duplex DNA as a third strand to form a triple helix. TFOs act in polypurine or polypyrimidine regions of DNA and bind DNA via Hoogsteen hydrogen bonds (Sargent et al. 2011, Pauwels et al. 2014).

Gene therapy remains a great promise for the treatment of genetic diseases. Recombinant AAV have been successfully used to transduce a variety of genes in different cellular types in vitro and have been validated in small and large animal models (in vivo). There are three critical elements in gene transfer: the gene, the target tissue and the vector. The gene is the active therapeutic agent, but the virus-derived vector is the determinant of the therapeutic success and of the toxicity profile (Potter et al. 2014). AAV vectors are currently being used in Phase I/II clinical trials for many diseases, such as CF, Pompe disease, α-1 antitrypsin deficiency, muscular dystrophy, Batten’s disease, Parkinson’s disease, Leber’s congenital amaurosis, hemophilia, and choroideremia (Mittermeyer et al. 2012, Zhong...
et al. 2012, Smith et al. 2013, MacLaren et al. 2014, Mendell et al. 2015). Furthermore, this treatment is already licensed in Europe for lipoprotein lipase deficiency (Mingozzi and High 2011).

More recently, the use of ZFNs has become a powerful strategy. These nucleases are engineered to introduce site-specific double-stranded DNA breaks in the genome. Site-specific alterations of the genome are then accompanied by homology-directed repair of the double-stranded DNA breaks using a linear double-stranded donor DNA fragment carrying the desired alteration. This strategy has been shown to be highly efficient at disrupting genes in several cell types and model organisms, making it a good choice for novel therapeutic applications (Granja et al. 2014). Although this approach can be remarkably effective, one obstacle of this approach is that ZFNs need to be designed and constructed for every specific location to be modified. Thus, ZFNs are most useful for repairing frequently occurring mutations (Aarts and Riele 2011).

A new approach that combines ZFNs with AAV has been studied. Based on the enzyme's ability to create a site-specific DNA double-strand break, Händel et al. (2012) demonstrated that ZFN-encoding AAV expression vectors can be employed to induce large chromosomal deletions or to disrupt genes.

Current strategies combine different techniques for sequence-selective double-strand DNA targeting, including TFOs, synthetic hairpin polynucleotides, engineered zinc finger proteins and peptide nucleic acids. TFOs bind as a third strand in the major groove of dsDNA. In the classical ‘pyrimidine motif’, thymine/cytosine containing TFOs hybridize to the complementary adenine/guanine bases of the target in a parallel orientation via Hoogsteen base pairing (Hansen et al. 2009).

Many of these strategies reviewed here are commonly used for missense, nonsense and frameshift mutations.

**FRAMESHIFT MUTATIONS**

Frameshift mutations change the reading frame, thus inducing a completely different translation from the original one. These mutations often inactivate genes by producing truncated, nonfunctional proteins (Morita et al. 2011).

The use of oligonucleotides in targeted sequence conversion has been developed for introducing sequence alterations, including deletions, insertions, and base-substitutions, into genomic DNA (de Semir and Aran 2006, Parekh-Olmedo and Kmiec 2007). Studies have demonstrated that both single-stranded DNA (ssDNA) fragments containing the sense sequence and the tailed duplex (TD) DNA fragments, prepared by annealing an oligonucleotide to the ssDNA fragment, have the ability to correct single-base substitution mutations. However, the ssDNA fragments correct mutations with low efficiency. Because the correction efficiencies of single-base substitution mutations by the TD fragments are higher, it is important to consider and examine the ability of the TD fragments to correct frameshift mutations (Morita et al. 2011).

Another approach for frameshift mutations therapy is the use of engineered zinc-finger proteins to recognize a unique chromosomal site, which can be fused to a nuclease domain. Moreover, double-strand breaks induced by the resulting ZFN can create specific sequence alterations by stimulating homologous recombination between the chromosome and an extrachromosomal DNA donor (Urnov et al. 2005, Rahman et al. 2011). Urnov et al. (2005) showed that ZFNs designed against an X-linked severe combined immune deficiency (SCID) mutation in the IL2Rγ gene yielded higher than 18% of the gene-modified human cells, even without selection.

The targeted gene correction technique employs a site-specific DNA lesion to promote a homologous recombination that eliminates the mutation in a gene of interest. However, double-strand breaks that are typically used to initiate
corrections can also result in genomic instability if deleterious repair occurs rather than gene correction, possibly compromising the safety of the targeted gene correction (Davis and Maizels 2011).

Cell therapy combined with gene therapy is a broadly expanding field in the correction of frameshift mutations and other mutations. Using a transgenic/knockout sickle cell anemia mouse model containing the human βS-globin gene, Chang et al. (2006) prepared embryonic stem (ES) cells from blastocysts that had the sickle cell anemia genotype and carried out homologous recombination with DNA constructs containing the βS-globin gene. Hematopoietic cells differentiated from these ES cells, produced both hemoglobin A and hemoglobin S, having the potential to correct the sickle and β-thalassemia mutations.

The development of induced pluripotent stem (iPS) cells allowed stem cell therapies to advance to new frontiers. Using a humanized sickle cell anemia mouse model, Hanna et al. (2007) showed that mice can be rescued after transplantation with hematopoietic progenitors obtained in vitro from autologous iPS cells. This was achieved after correcting the human sickle hemoglobin allele with gene-specific targeting. Ye et al. (2009) treated fibroblasts from patients with β-thalassemia caused by frameshift mutations. They showed that iPS cells could be produced from the somatic cells of these patients and that the mutations could be corrected with gene targeting. Cells differentiated into hematopoietic cells can be returned to the patient. The innovation of reprogramming somatic cells into induced pluripotent stem cells provides many possible new approaches for treating β-thalassemia and other genetic diseases.

Nature provides an interesting model of mutation reversion that should be thoroughly explored. Revertant mosaicism is a naturally occurring phenomenon that involves the spontaneous correction of a pathogenic mutation in a somatic cell. Recent studies suggest that this is not a rare event and that reversion could be clinically relevant to phenotypic expression and patient treatment. Indeed, revertant cell therapy represents a potential ‘natural gene therapy’ because in vivo reversion obviates the need for further genetic correction. Revertant mosaicism has been observed in several inherited conditions, including epidermolysis bullosa, a heterogeneous group of blistering skin disorders (Lai-Cheong et al. 2011).

Several structural modifications and techniques are being developed to optimize the therapeutic responses to these new treatments. However, it is also apparent that much remains to be learned regarding the organization of these modifications. Eventually, principles with predictive value should emerge to guide the use of these new approaches.

CONCLUSIONS

Translational research for rare diseases is clearly a resource-intensive undertaking in terms of cost and time. Therefore, the central aspects of any broadly successful approach for this class of diseases will be to create and employ generalizable methodologies whenever possible. The identification of novel, non-toxic and efficient therapeutic agents and advances in the stability and delivery of antisense oligonucleotide splice-correction therapies are necessary. Although the approaches reviewed in this paper are credible and feasible, the prospect of a rapid configuration of numerous effective disease therapies should be viewed in perspective. Among the many genetic disorders that have been or will shortly be molecularly characterized, only a small number will be tractable with the approaches reviewed here; however, the generation of information will at least help to move a larger number of disorders closer to the day of effective therapy.

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**RESUMO**

Com o avanço e a popularização de técnicas moleculares, a identificação de mutações genéticas que causam doenças aumentou drásticamente. Deste modo, o número de laboratórios disponíveis para investigar uma determinada doença e, consequentemente, o número de diagnósticos, aumentou ao longo do tempo. Embora seja necessário identificar mutações e fornecer diagnósticos, é crucial também desenvolver abordagens terapêuticas específicas baseadas nestas informações. Esta revisão tem como objetivo destacar os avanços recentes em terapias mutação-alvo com produtos químicos que mitiguem a patologia mutacional ao nível molecular, para doenças que, para a maior parte, não têm tratamento eficaz. Atualmente, existem várias estratégias sendo utilizadas para corrigir diferentes tipos de mutações, incluindo as seguintes: a identificação e caracterização de compostos *readthrough* tradicionais; redirecionamento de *splicing* mediado por oligonucleotídeos de sentido contrário; reparo por *mismatch*; e salto de exon. Estas terapias e outras abordagens são revistas neste artigo.

**Palavras-chave:** redirecionamento *antisense* de *splicing* mediado por oligonucleotídeo, remoção de exon, reparo por *mismatch*, terapias alvo-molecular, *readthrough* tradicional.

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MITIGATING MOLECULAR PATHOLOGY USING CHEMICALS


TAYLOR JK, ZHANG QQ, WYATT JR AND DEAN NM. 1999. Induction of endogenous Bel-xS through the control of...


