Review

Succinic Semialdehyde Dehydrogenase Deficiency: An Update

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Abstract: Succinic semialdehyde dehydrogenase deficiency (SSADH-D) is a genetic disorder that results from the aberrant metabolism of the neurotransmitter γ-amino butyric acid (GABA). The disease is caused by the impaired activity of the mitochondrial enzyme succinic semialdehyde dehydrogenase. SSADH-D manifests as varying degree of mental retardation, autism, ataxia and epileptic seizures, but the clinical picture is highly heterogeneous. So far, there is no approved therapy for this disease. In this review, we briefly summarize the molecular genetics of SSADH-D, the past and ongoing clinical trials and the emerging features of the molecular pathogenesis, including redox imbalance and mitochondrial dysfunction. The main aim of this review is to discuss the potential use of further therapy approaches that have so far not been tested in SSADH-D, such as pharmacological chaperones, read-through drugs and gene therapy. Special attention will also be paid to elucidating the role of patient advocacy organizations in facilitating research and in the communication between the researchers and the patients.

Keywords: Succinic semialdehyde dehydrogenase deficiency; gamma-amino butyric acid; organic acidurias; enzyme replacement therapy; pharmacological chaperones; clinical trials; autophagy

1. Succinic Semialdehyde Dehydrogenase Deficiency: Clinical Phenotype, Genetics and Standard Care

1.1. Clinical Phenotype and Diagnosis of Succinic Semialdehyde Dehydrogenase Deficiency

Succinic semialdehyde dehydrogenase deficiency (SSADH-D, also called 4-hydroxybutyric aciduria, OMIM #271980) is an ultra-rare monogenic disorder of the γ-amino butyric acid (GABA) metabolism, with approximately 400 patients known to literature (reviewed in [1]). The first case of SSADH-D was identified in 1981 in a patient excreting γ-hydroxybutyric acid (GHB) in the urine, and only two years later, the deficiency of succinic semialdehyde dehydrogenase (SSADH) activity was demonstrated to be the underlying cause [2, 3]. Variants of the ALDH5A1 gene have later been shown to be the cause of SSADH-D [4]. SSADH-D is inherited in an autosomal recessive fashion, and both parents typically present as carriers. Heterozygous carriers of one defective allele show no clinical signs of the disease, whereas the patients who are either homozygous or compound heterozygous for disease-causing variants are affected to a varying degree. Enzymatic dysfunction of SSADH leads to an accumulation of potentially neurotoxic metabolites, including GABA and GHB. Despite ambitious scientific effort, detailed knowledge about many aspects of the pathophysiology
of the underlying enzyme defect is still lacking, and at present, no curative treatment is available for SSADH-D. As with many other rare disorders affecting the central nervous system (CNS), several symptomatic treatments have been and are currently being investigated [5, 6].

The clinical picture of SSADH-D is highly heterogeneous, and in many cases, the somewhat nonspecific nature of the symptoms may delay the diagnosis of patients without prior family history of the disease [1, 7]. However, common manifestations of SSADH-D include a varying degree of mental retardation, psychiatric disorders, autism-like symptoms and impaired speech, along with sleep disturbances [1, 8, 9, 10]. Some degree of developmental delay and intellectual disability are found in all patients, while around 80% of the patients are affected by ataxia and muscular hypotonia [7]. Starting in late childhood, most patients (around 60%) develop epileptic seizures, ranging from absence seizures to generalized forms of epilepsy, which are also present in adult patients. Apart from the seizure phenotype, the disease usually does not exhibit a further progressive course. However, the highly variable clinical presentation of SSADH-D frequently makes diagnosis difficult. Already in 2004, Gordon et al. have drawn attention to the very poor genotype/phenotype correlation in SSADH-D [11]. Even in a single family with two or more affected children harboring the same pathogenic variants of SSADH, the degree of disability and the symptoms can vary greatly [12]. The exact cause of this variability despite the same genetic background is not known. Currently, active research efforts are pursued to better understand the causative relationship between the molecular defect and the subsequent clinical consequences. In addition, the detailed molecular consequences of specific SSADH disease-causing variants in terms of SSADH enzyme function are actively characterized.

SSADH-D is caused by a defect in the catabolism of GABA, the main inhibitory neurotransmitter of the CNS. Excess GABA is usually removed by the successive action of enzymes mediating its degradation (Figure 1), such as GABA transaminase that removes the amino group in GABA, producing succinic semialdehyde (SSA). This metabolic product is in turn removed by the SSADH enzyme that converts it into succinic acid, which can be further metabolized in the tricarboxylic acid cycle. In SSADH deficient patients, the GABA metabolic pathway is disrupted due to low or absent activity of SSADH. Due to this, SSA cannot be processed by its normal catabolic pathway, resulting in accumulation of GHB that is generated from SSA by an aldo-keto-reductase. In SSADH-D, a high degree of accumulation of both GABA and GHB is observed in the CNS as well as in peripheral tissues and body fluids of the patients [11]. However, it is not clear which one of these neuroactive substances is the main contributor to the neuronal impairment observed in SSADH-D, since large excess of either of these compounds can exert neurotoxic effects. In fact, GHB is also known as a “rape drug”, as it can be misused to induce amnesia, which has led to considerable forensic interest towards GHB. In the nervous system, GHB can act by binding to its own receptor, but it has been shown that GHB also binds to certain GABA_\alpha receptors [13], whereas GABA itself acts on both GABA_\alpha and GABA_\gamma receptors [14]. Thus, GABA receptors have been a target of research aiming at clinical interventions in SSADH-D by applying receptor agonists or antagonists. It is out of scope of this review to discuss the function of GABA as neurotransmitter, and we would like to recommend the reviews by Petroff and Siucinska to interested readers [15, 16]. However, clinical and pre-clinical interventions of GABA and GHB receptors in SSADH-D will be briefly discussed in the present review.

The diagnosis of SSADH-D is presently based on detection of GHB in the organic acid profile in urine [11]. In addition, GHB can be measured in other body fluids [17, 18], but this is usually done only for research purposes and is not routinely used. In addition, genetic testing can be the first diagnostic step and should always follow a biochemical suspicion for confirmation of the diagnosis. Since SSADH-D can present as an autism spectrum disorder [19], the ALDH5A1 gene is included in the commercially available autism gene panel tests [20]. Applying such panels to autism cases of unknown etiology could thus help to identify new cases of SSADH-D. A large number of cases is still suspected to remain undiagnosed, considering the inhomogeneous global distribution of diagnosed cases [21]. The diagnostic rate could also be improved e.g. by introducing newborn screening
programs for organic acidurias, since it has recently been shown that diagnosis for SSADH-D may be possible from a dried bloodspot [22].

Figure 1. Overview of GABA catabolism. Neuronal GABA synthesis is mainly accomplished from glutamine derived from astrocytes and from protein and fat catabolism. Glutaminase converts glutamine to glutamate, followed by glutamate decarboxylase that converts glutamate to γ-aminobutyric acid (GABA), the most abundant inhibitory neurotransmitter of the CNS. Degradation of GABA is accomplished by GABA transaminase that catalyzes the transamination of GABA to succinic semialdehyde (SSA), which in turn is converted to succinate by SSADH. Dysfunction of SSADH causes accumulation of SSA and GABA and subsequent conversion of these metabolites to γ-hydroxybutyric acid (GHB) through SSA reductase.

1.2. Standard care and emerging picture of the pathophysiology in SSADH-D

There is no curative treatment for SSADH-D to date, and the patients are thus treated mainly symptomatically. This is highly unsatisfactory for both patients and clinicians, and novel therapeutic approaches for treating the actual cause of the disease are needed. Conventional treatment approaches include speech, physio and occupational therapy, whereas with older patients, psychotherapy is also frequently applied. One major treatment goal is the control of epileptic seizures in those patients where they occur [8]. A broad spectrum of antiepileptic medications has been used in patients with SSADH-D, including the sodium channel blockers lamotrigine and carbamazepine [5], whereas valproate should be avoided due to its inhibition of potential residual enzymatic activity [23]. For vigabatrin, which is an irreversible inhibitor of GABA transaminase and thus inhibits the subsequent formation of SSA, inconsistent results and severe side effects have been shown (see section 3.2 and [24, 25]). In addition to these drugs, medications against a broad spectrum of symptoms are in use in SSADH-D, such as antidepressants and substances against anxiety and inattention [9, 26]. However, all these agents are designed to act only against specific symptoms and do not address the actual pathomechanisms.

Due to the role of SSADH in GABA catabolism, it was originally suggested that the clinical phenotype is mainly based on a fundamental imbalance in the amount of GABA, the major inhibitory transmitter of the CNS, and the toxic GHB. In fact, both substances can be found in elevated levels in white and grey brain matter as well as in urine and plasma of SSADH-D patients [27]. Recently, it has also become evident that further molecular events may damage specific cellular functions and thus contribute to the clinical picture. For example, in vitro experiments and the murine model have pointed to a redox imbalance, mitochondrial dysfunction and altered signaling through the mTOR (mechanistic target of Rapamycin) pathway in SSADH-D. In addition, alterations in the expression
of numerous genes have been shown, which might contribute to the phenotype e.g. by affecting ion efflux/influx in neuronal cells. In Chapter 3, we will summarize these findings and elucidate on their potential therapeutic relevance in SSADH-D.

2. The ALDH5A1 gene and the SSADH enzyme: lessons learned from disease models

2.1 ALDH5A1 gene splicing isoforms and genetic variants in SSADH-D

The ALDH5A1 gene resides in the human chromosomal region 6p22 [4, 28], and the gene structure has been characterized almost twenty years ago [29]. According to databases such as GenBank, three human isoforms that show differences in the coding region have been predicted to exist. These isoforms arise due to alternative splicing of two in-frame exons in the pre-mRNA, and the isoforms 1 (NM_170740.1) and 2 (NM_001080.3) differ from each other in length by 39 bases (13 amino acids), with isoform 1 being the longer one. Isoform 3 (NM_001368954.1) has been predicted to lack further 144 bases (48 amino acids) within the coding region, and it thus represents the shortest of the three isoforms. Figure 2 shows the predicted protein sequences of the three isoforms. According to our experience, isoform 2 mRNA appears to be the major form that is present in most cells, whereas we have found only trace amounts of isoform 1 mRNA, as detected by quantitative real-time PCR (M.D., A.B. and R.T., unpublished findings). To our best knowledge, no experimental evidence has been put forward that supports the existence of the predicted splicing isoform 3, neither at mRNA nor at protein level. The additional 13 amino acids in the isoform 1 translation product have been postulated to result in an inactive SSADH enzyme [29], and the same is likely to be the case for isoform 3, in which 48 amino acids would be missing from the cofactor binding domain of SSADH [30]. Thus, isoform 2 should be considered as the major enzymatically active form of SSADH in humans.

Various types of gene variants including missense, nonsense and splicing mutations, deletions and insertions in the ALDH5A1 gene have been shown to be responsible for SSADH-D (see section 2.3 and e.g. [31, 32]). By 2016, as many as 45 pathogenic variants were described, and further mutations are identified as more patients receive a correct diagnosis [24]. In many families, homozygous gene defects are found due to consanguinity of the parents. In addition to known pathogenic variants, various single nucleotide polymorphisms (SNPs) of unknown significance have been described in the respective databases (see e.g. [33]).

Unfortunately, major confusion can be caused upon diagnosis by the use of different gene isoforms for the numbering of the detected variants. Many diagnostic companies base their numbering on the isoform 1, which is not the major splicing isoform of SSADH. Therefore, the numbering of all amino acids beyond Glu242 (the site of insertion of the additional 13 amino acids) is shifted by 13, and care has to be taken when analyzing the consequences at the amino acid level. In this review, the numbering of the genetic and protein variants is based on the isoform 2 and the protein translated from it.

2.2. SSADH enzyme function and structure

The SSADH enzyme precursor polypeptide of 535 amino acids (isoform 2) is synthesized in the cytosolic, free ribosomes. This polypeptide contains a predicted mitochondrial targeting sequence (MTS) of 47 amino acids in its amino terminus (Figure 3). This MTS has been suggested to be responsible for the mitochondrial matrix targeting of the SSADH precursor, although experimental evidence has so far not been presented. In the mitochondrial matrix, SSADH folds and forms oligomers, most likely tetramers, that represent the enzymatically active form [34].

The 3-dimensional structure of the recombinant human SSADH enzyme (amino acids 48-535) has been solved with a resolution of 2.0 Å [34]. Each asymmetric unit of the crystal structure contained the monomeric form of the polypeptide, but a tetrameric structure was predicted for SSADH by applying the crystallographic F432 symmetry model. Three domains that are responsible for the cofactor NAD⁺ binding (amino acids 48-173, 196-307 and 509-524), for the catalysis (amino acids 308-508) and for the tetramerization (amino acids 174-195 and 525-535) can be identified in each
monomeric structure ([34], see also Figure 3). The first two domains mainly exhibit α-helical and β-sheet structures, whereas the tetramerization domain contains three β-strands in antiparallel orientation.

![Figure 2](image-url)

**Figure 2. SSADH isoform protein sequences.** Numbers to the left in bold show the isoform numbers. Blue highlight marks the 13 extra amino acids in isoform 1, whereas green highlight depicts the 48 amino acids present in isoforms 1 and 2 but missing from isoform 3.

![Figure 3](image-url)

**Figure 3. Exon and domain structure of SSADH, and localization of selected SSADH-D missense variants.** The human SSADH isoform 2 exhibits 535 amino acids. Green boxes show the exons, joined by the black lines representing introns, the length of which is given above the lines. Colored blocks refer to the protein domains as indicated. Numbers below the bar refer to the first amino acid of the respective domain. Dashed lines connect the selected disease-causing variants with the respective exons and protein domains.
SSADH is a member of the aldehyde hydrolase (ALDH) protein family, and its 3D structure shows a clear homology to other known members of this family. However, the active site and the substrate entrance exhibit features that are specific to SSADH. This is consistent with the fact that, as compared to other members of the ALDH family, SSADH shows a remarkable substrate specificity towards SSA. Highly intriguingly, Kim and coworkers showed that the accessibility of the SSADH active site appears to be mediated by a redox-switch mechanism, which may explain the high substrate specificity of SSADH [34]. In the crystal structure of SSADH that was obtained in the absence of a reducing agent, the catalytic Cys340 within the active site appears to be engaged in a disulfide bond with Cys342. This results in a conformation that would prevent the entry of both the substrate SSA and the cofactor NAD⁺ to their respective binding sites, resulting in a catalytically inactive state of the enzyme. Exchange of the Cys342 residue prevented the formation of this closed conformation and resulted in a higher enzyme activity. The authors suggested that the activity of human SSADH might physiologically be controlled by oxidative stress and the redox state within the mitochondrial matrix. This is especially intriguing since changes in the redox status have been shown to be a feature of SSADH-D (reviewed in [30]), and these alterations might also contribute to the profound lack of residual activity that has been observed in the case of most of the disease-causing variants.

### 2.3 Consequences of the pathogenic variants found in SSADH-D

Residual SSADH enzyme activity of pathogenic ALDH5A1 variants has been investigated in several studies based mainly on overexpression of the variants (see e.g. [31, 32, 35, 36, 37, 38]). However, so far, it has not been possible to conclusively establish how each individual variant affects the enzyme activity in patients. As the residual enzyme activities may vary largely even among siblings with the same genotype, it is likely that further factors play a major role in regulating to which degree the genetic variants impair the SSADH enzyme activity and function. Identifying these factors would be of major interest, as this might provide novel means of ameliorating the consequences of the lack of SSADH activity and facilitate the generation of novel therapies for SSADH-D. With technological progress that facilitates a more detailed analysis, more attention should be paid to genotype/phenotype correlation, and both experimental and in-silico approaches such as modeling the structure of the pathogenic variants should be used to elucidate how a specific genetic defect affects the function of the SSADH enzyme.

Numerous pathogenic variants localized throughout the SSADH polypeptide have been identified in SSADH-D patients. Most of these variants affect the predicted cofactor binding domain, but this does not necessarily imply that this domain is more vulnerable than other regions of SSADH. It may rather reflect the fact that the NAD-binding domain occupies almost half of the enzyme, with 254 of the 535 amino acids being part of this domain (see Figure 3). The structural and functional consequences of some of the disease variants have been analyzed, but most of them appear to have a very profound effect on the SSADH enzyme activity, at least when analyzed in an overexpression system (see e.g. [29, 31]). Table 1 summarizes some of the predicted and experimentally addressed consequences of SSADH-D missense variants (see also Figure 3).

Based on the crystal structure of SSADH, Kim and colleagues suggested that many of the missense variants that contain substitutions in an amino acid buried deep within the polypeptide (e.g. Cys93Phe, Cys223Tyr, Thr233Met and Pro382Leu) could result in a profound destabilization of the enzyme structure [34]. Many other published variants (e.g. Gly176Arg, Gly176Glu, Pro182Leu and Gly533Arg) are localized within the tetramerization domain [29, 31, 39], and the replacement of the small Gly residues by bulky and charged residues such as Arg or Glu is highly likely to impair SSADH oligomerization and thus also its stability. Consistent with these predictions, our unpublished findings in patient fibroblasts exhibiting some of these variants show that the amount of the SSADH polypeptide detected by Western Blot is highly reduced. Some SSADH-D variants may also affect substrate or cofactor binding and thus result in the impairment of the catalytic activity. For example, Gly268 is involved in the binding of the cofactor NAD⁺. Thus, the SSADH-D variant with a
substitution Gly268Glu is likely to impair NAD⁺ binding and thus exhibits a strong negative effect on the catalytic activity of SSADH [34].

**Table 1.** Selected missense variants of the human *ALDH5A1* gene that have been associated with SSADH-D. Numbering of the genetic variants is based on the human isoform 2 (NM_001080.3).

| Genetic variant | Amino acid change | Domain | Remark, severity | Ref |
|----------------|-------------------|--------|-----------------|-----|
| c.106G>C       | Gly36Arg          | MTS    | Mild            | [29, 31] |
| c.278G>T       | Cys93Phe          | NADB   | Several families, severe | [31] |
| c.526G>A       | Gly176Arg         | Oligom.| Severe, conserved residue | [32] |
| c.527G>A       | Gly176Glu         | Oligom.| Conserved residue | [39] |
| c.538C>T       | His180Tyr         | Oligom.| Mild, exacerbating | [29, 31, 32] |
| c.545C>T       | Pro182Leu         | Oligom.| Mild, exacerbating | [29, 31] |
| c.691G>A       | Glu231Lys         | NADB   | Conserved residue | [39] |
| c.668G>A       | Cys223Tyr         | NADB   | Severe           | [31] |
| c.709G>T       | Ala237Ser         | NADB   | Mild, exacerbating? | [29, 31, 32] |
| c.698C>T       | Thr233Met         | NADB   | Severe           | [31] |
| c.763A>G       | Asn255Asp         | NADB   | Several families, severe | [31] |
| c.764A>G       | Asn255Ser         | NADB   | Intermediate     | [31] |
| c.800T>G       | Val267Gly         | NADB   | Conserved residue | [39] |
| c.803G>A       | Gly268Glu         | NADB   | Cofactor binding, severe | [31, 34] |
| c.901A>G       | Lys301Glu         | NADB   | Cofactor binding | [40] |
| c.1005C>A      | Asn335Lys         | Catal. | Dynamic loop, severe | [31, 34] |
| c.1115A>G      | Asn372Ser         | Catal. | Not characterized | [29] |
| c.1145C>T      | Pro382Leu         | Catal. | Severe           | [31] |
| c.1145C>A      | Pro382Gln         | Catal. | Not characterized | [31] |
| c.1126G>A      | Val406Ile         | Catal. | Not characterized | [29] |
| c.1226G>A      | Gly409Asp         | Catal. | Severe           | [31, 38] |
| c.1267A>T      | Thr423Ser         | Catal. | Mild, exacerbating? | [32] |
| c.1498G>C      | Val500Leu         | Catal. | Substrate binding, severe | [38] |
| c.1529C>T      | Ser510Phe         | NADB   | Not characterized | [39] |
| c.1597G>A      | Gly533Arg         | Oligom.| Severe           | [31] |

1 MTS = mitochondrial targeting sequence; NADB = NAD binding domain; Oligom. = oligomerization domain; Catal. = catalytic domain; 2 Severity of the mutation refers to its effect on the SSADH enzyme activity in overexpression systems.

In addition to the known SSADH-D variants, various SNPs that result in an amino acid exchange have been detected in the coding region of SSADH. For most of these variants, the significance and relevance for the enzyme function and activity is unknown. One exception is the His180Tyr variant (SNP rs2760118) whose consequences have been analyzed in an overexpression system [29, 31]. When overexpressed, the His180Tyr SSADH mutant has 83% of wildtype activity, and should thus have no profound pathogenic relevance. However, when His180Tyr is expressed in combination (as a double mutation in the same construct or in the same allele in the case of patients) with another mild SSADH variant such as Pro182Leu, it appears to reduce the residual enzyme activity of its counterpart even further [31]. Interestingly, His180 and Pro182 are localized in the oligomerization domain, so they may cause only a slight impairment of SSADH tetramerization when present alone. However, they appear to produce a more profound effect when combined with another SSADH variant. Therefore, despite their mild individual effect on SSADH enzyme activity, His180Tyr and Pro182Leu substitutions cannot be considered as harmless or completely non-pathogenic, and their relevance needs to be considered in the context of other substitutions that are present in the same allele. Unfortunately, according to our experience, the SNPs resulting in amino acid exchanges that are listed as non-pathogenic in the databases are frequently ignored and not disclosed by companies.
providing genetic diagnosis for SSADH-D. However, for the researchers and clinicians, it would be essential to know the exact genotypic variants present in a patient, in order to be able to correctly dissect the potential molecular consequences of the mutations. Thus, all sequence variants of the mutated gene should always be disclosed in the diagnosis files, as it is likely that many patients will have additional substitutions such as His180Tyr, which has been shown to be present in more than 30% of analyzed chromosomes within the GnomAD exome database [41].

Most of the studies addressing the consequences of SSADH-D gene variants have measured the residual enzyme activity only, without paying attention to the expression level, mRNA amount, or localization of the enzyme. Furthermore, much of the data are derived from overexpression systems. Therefore, in the future, it will be important to perform such studies in more suitable systems, especially in patient-derived cells, such as fibroblasts, or neuronal cells derived from induced pluripotent stem cell lines (see below). Furthermore, when using overexpression studies, the effect of specific variants should always be characterized in cells that lack endogenous SSADH. This is important since SSADH is a tetramer, and oligomerization of the mutant enzyme with the endogenous wildtype SSADH might not reveal the true degree of functional impairment of the variant, as compared to the expression of the variant alone. We have recently generated CRISPR-based knockout HEK293T cells that do not express endogenous SSADH, which will be a valuable tool to analyze the molecular consequences of SSADH mutations in a context without endogenous background (M.D., A.B. and R.T., unpublished data).

2.4 Disease models: Cellular models, organoids and SSADH knockout mouse

The simplest models for studying SSADH-D are cell culture-based systems, either ectopic overexpression of the enzyme in a suitable cell line such as HEK293 cells, or patient cells, e.g. dermal fibroblasts that can easily be obtained and cultured. The advantages of these systems are the ease of use and low price of culture reagents. However, overexpression may result in aberrant behavior of the protein due to aggregation or saturation of machineries required for the mitochondrial translocation. Dermal fibroblasts from a patient express low levels of SSADH, but they exactly represent the genotype of the patient. On the other hand, their life span is very limited, and finding a suitable isogenic control may be difficult. Therefore, modern genome editing methods such as the CRISPR/Cas technology can be used to produce genomic SSADH variants that mimic the patient genotype in a suitable cell line.

Patient-derived fibroblasts or peripheral blood mononuclear cells (PBMCs) can be reprogrammed to induced pluripotent stem cells (iPSCs). Great advances have been made in this technology in recent years, and it is becoming ever more popular as a model system to study human diseases. The advantage of this model is that iPSCs can be differentiated into all cell types of the body, especially those that represent the tissue of interest, such as the liver or the brain. This can be done in a 2-dimensional culture in a dish, and established protocols are available for various cell types. Although differentiated cells, such as neurons, representing the patient genotype are highly desirable, the generation, culture and differentiation of iPSCs is not trivial and requires expensive cell culture media and supplements. Recently, diverse protocols for the differentiation of iPSCs in 3D culture systems to so-called organoids have been published. These include protocols for the generation of cerebral organoids [42], but also regionalized brain organoids like midbrain or forebrain organoids [43, 44, 45]. These organoids usually consist of several cell types organized in structures resembling the organs in question. This method is highly sophisticated, and the organoids recapitulate the situation in patients to a higher degree than 2D cultures. However, the method requires special skills, very expensive reagents, and the organoid development usually takes several weeks. Therefore, organoids will surely be very useful in the future for studying diseases and testing therapies, but they are unlikely to replace the 2D culture systems completely. Moreover, they are not easy to use for high content screening approaches due to their high batch to batch variations and the difficulties in finding a suitable readout for screening purposes.

The murine SSADH knockout (Aldh5a1−/−) model was developed almost two decades ago by Hogema and colleagues [46]. These mice are well accepted as an SSADH-D animal model, although
the phenotype is far more severe than what is seen in most human SSADH-D patients. The characteristics of these mice have recently been very thoroughly summarized by Kim et al. [30]. We thus kindly refer the readers to this excellent review, and we here only elaborate the major features of this mouse model. The SSADH knockout mice develop generalized seizures by 3-4 weeks of life, resulting in an early death. The metabolic features associated with these mice well recapitulate those of the human SSADH-D patients, e.g. elevated GABA and GHB levels in tissues and in the urine and absence seizures during the second week of life, which tend to culminate into more generalized seizures in the third week of life. Many of the findings observed in SSADH-D patients are found in this mouse model in an exacerbated form. Therefore, the SSADH knockout model currently represents the best available animal model that mimics the main features associated with the human disease.

For potential treatment trials in the animal model, the early lethality poses a major problem as any treatment would have to be started very early (latest around day 10), where the pups are still quite small in size and may be difficult to handle. On the other hand, the extended survival beyond the expected lifetime of 3-4 weeks can be used to monitor the treatment effect. However, treatment trials to monitor long-term benefits are not possible. Therefore, an animal model that more precisely mimics the mainly slow course of the disease in humans would be highly desirable, and could be obtained by transgenic or CRISPR/Cas-mediated knock-in of a suitable disease-causing genetic variant.

2.5. Mitochondrial dysfunction, redox imbalance and autophagy defects in SSADH-D

The lack of SSADH enzyme activity results in alterations in the amount of GABA and GHB that are likely to be directly relevant for the pathogenesis and the symptoms observed. However, indirect effects that result in organelle dysfunction or impairment of signaling have been suggested to contribute to SSADH-D. For example, oxidative damage has been shown to be a prominent feature in both patients and in the murine SSADH-D model, and mitochondrial aberrations are frequently observed ([47] and reviewed in [30]). Interestingly, GHB has been shown to inhibit lipid biosynthesis and to induce oxidative stress in rats [48, 49]. Silva and colleagues suggested that altered lipid amounts were not caused by a direct inhibition of the enzymes involved in lipid biosynthesis by GHB, whereas mitochondria were likely to contribute to this effect [49]. These findings again point to the importance of indirect toxic effects in the disease pathogenesis of SSADH-D.

A subset of SSADH-D patients shows rather non-specific signal changes in their brain magnetic resonance imaging (MRI). Signs of metabolic toxicity can be revealed by MRI of patient brains [7, 50], pointing to an important contribution of oxidative damage. This is consistent with the findings in other organic acidemias such as methylmalonic acidemia [51, 52], showing that redox imbalance and mitochondrial dysfunction are a typical feature also in these diseases. Beyond that, aberrations in lipid biosynthesis have also been shown in organic acidemias [53], which may lead to aberrant brain development and myelination already before birth and postnatally. In the developing brain, lipids such as gangliosides are of vital importance for the proper brain maturation and neurite outgrowth [54, 55]. In the SSADH knockout mouse model, downregulation of several genes associated with myelination and reduced levels of major myelin plasmalogen components have been observed [56]. Thus, impairment of lipid homeostasis and myelination may also be a common feature in SSADH-D and other organic acidemias.

Several studies have addressed the role of oxidative stress/damage in SSADH-D. Using the mouse model, Latini and colleagues showed that the total radical-trapping antioxidative potential (TRAP) and glutathione (GSH), which represent the non-enzymatic antioxidative defense mechanisms in tissues, were highly reduced in some tissues, especially in the liver and in cerebral cortex [57]. Low GSH levels have also been observed in an SSADH-D patient, who also exhibited high organic acid derivate levels in the urine [47]. Latini and coworkers also observed an increased lipid peroxidation in the cerebral cortex and the liver of the SSADH knockout mice, demonstrating that oxidative damage to lipids takes place in these tissues [57]. Interestingly, SSADH has been shown to be the major enzyme that is required for the elimination of 4-hydroxy-trans-nonenal (HNE), a lipid...
peroxidation product, in the brain [58]. HNE has also been shown to be involved in the induction of oxidative stress in other diseases such as Alzheimer’s disease [59, 60]. In SSADH-D, it is thus likely that in the absence of SSADH activity, HNE may not be disposed of in a proper way, and its increased levels may further contribute to oxidative stress.

Lakhani and coworkers have shown that in yeast, GABA is involved in the regulation of specific forms of autophagy, mitophagy and pexophagy, by increasing the activation of the mTOR complex [60]. Autophagy is an important clearance process during which altered cell organelles such as mitochondria (mitophagy) and peroxisomes (pexophagy) are eliminated via lysosomal uptake and degradation (for review, see [61, 62]). Impaired mitophagy and pexophagy as such can result in increased production of reactive oxygen species (ROS) and oxidative stress, which are important pathogenic features in SSADH-D. In line with this, SSADH deficient mice exhibit increased numbers of mitochondria that show morphological defects, which would be consistent with impaired mitophagy [60]. Defects in autophagy are also associated with diverse neurodegenerative disorders, including the neuronal ceroid lipofuscinoses, Alzheimer’s disease and Parkinson’s disease, which also show mitochondrial dysfunction and increased oxidative stress [61, 63].

The activation state of the mTOR complex 1 (mTORC1) is a key element in the regulation of autophagy. The active mTORC1 is associated with lysosomes where it inhibits two major inducers of autophagy, the Atg1 kinase and the Atg13 protein. Nutrient starvation is associated with dissociation of the inactive mTORC1 from the lysosomes, resulting in increased autophagy. Thus, activation of mTOR by GABA is likely to exacerbate the accumulation of aberrant mitochondria in SSADH-D by inhibiting mitophagy, suggesting that mTOR inhibition may exhibit therapeutic potential in SSADH-D. Vogel and coworkers have analyzed the effect of various mTORC inhibitors in the SSADH knockout mice [64, 65]. They showed that the lifespan of the mice can be extended by dual inhibitors of mTORC1 and mTORC2, Torin 2 and XL-765 [65]. In addition, the mTOR-independent autophagy inducing peptide tat-Beclin 1 [66] also prevented the early lethality in these mice [65]. These findings suggest that inhibition of mTOR activity and/or induction of autophagy may exhibit therapeutic potential in SSADH-D.

Modulation of autophagy is expected to show high potential for diseases as diverse as metabolic disorders, cancers, neurodegenerative and infectious diseases [67, 68], and drugs that modulate the autophagic flux and exhibit therapeutic potential for various diseases have been identified in drug screens [68]. Disease-specific autophagy drug screens may also reveal novel pathways to be targeted for autophagy induction, as has recently been shown for juvenile neuronal ceroid lipofuscinoses, where pathways such as calcium signaling, isoprenoid pathway and microtubule dynamics were identified as potential drug targets for ameliorating autophagy defects [69]. Therefore, drug screens for further potential autophagy modulators could be carried out also for SSADH-D to identify novel disease-specific pathways that could be targeted to enhance autophagy.

3. Therapy Options for SSADH Deficiency

3.1 Current and past clinical trials in SSADH-D

For three decades, scientists have strived to find a pharmaceutical approach to compensate for the problems caused by a dysfunctional SSADH enzyme, e.g. by influencing the levels of unwanted by-products that stem from the oversupply of GABA. Ideas for curative treatment have also started to emerge, as our understanding of the molecular pathology of the disease is increasing. Some clinical trials targeting the receptors involved in GABA and GHB function have been carried out in the recent years, with varying success. These treatment concepts and previous clinical trials are summarized in the following paragraphs and in Table 2. Since the treatments and clinical trials have recently been excellently addressed by Vogel and colleagues, we here only shortly summarize the trials and the rationale of these treatments [70].
Table 2. Summary of the preclinical treatment concepts and clinical trials that have been carried out or are in progress for SSADH-D.

| Intervention | Primary target | Mode of action | Outcome in preclinical models | Clinical trial and outcome |
|--------------|----------------|----------------|-------------------------------|---------------------------|
| SGS-742 [71] | GABA<sub>B</sub> receptor | Antagonism | Improvement of epileptiform activity, reduced absence seizures in aldh5a<sup>−/−</sup> mice | Completed, phase 2, 19 patients enrolled [73] |
| CGP-35348 [72] | Diffuse GABA<sub>A/B</sub> modulatory receptor effects | Resuscitative Effect of an Antagonist | Improves survival of aldh5a<sup>−/−</sup> mice [75] | 1 patient, reversal of MRI-documented lesions [76], no effect in TMS [77] |
| Taurine [74] | GABA receptor | Antagonism | Improves survival of aldh5a<sup>−/−</sup> mice [81], MDCK cells [79] | - |
| NCS-382 [78, 79, 80] | GHB receptor | Antagonism | Improves survival of aldh5a<sup>−/−</sup> mice [46] | Effective in 1/3 of patients. Side effect: loss of vision [6] |
| Vigabatrin | GABA transaminase | Inhibition | Improves survival of aldh5a<sup>−/−</sup> mice | Increased level of GHB in urine with valproic acid [23] |
| Valproic acid [23, 82] | SSADH | Inhibition | - | - |
| Rapamycin, Torin, XL-765 [60, 64, 65, 83] | mTORC1/2 | mTORC inhibition, induction of autophagy | Improves survival of aldh5a<sup>−/−</sup> mice | - |
| Tat-Bec1 [83] | Beclin 1 | Beclin 1 independent induction of autophagy | Improves survival of aldh5a<sup>−/−</sup> mice, induces modest weight gain | - |
| Ketogenic diet [84, 85] | Neuroprotective effects | Neuroprotective effects | Improves survival of aldh5a<sup>−/−</sup> mice [84, 85] | - |

3.2. Clinical trials targeting the neurotransmitter receptors

The chemical hallmark of SSADH-D is an increased concentration of GABA and GHB in body fluids, such as blood, urine, and cerebrospinal fluid. It is therefore reasonable that the majority of the therapeutic trials have focused on substances that exhibit a direct effect on GABA and GHB receptor subtypes. Preclinical data also exist for a variety of further treatment approaches. These data were mainly generated using the Aldh5a<sup>−/−</sup> mouse model that exhibits an early lethal phenotype due to convulsive status epilepticus [46, 75]. The translation of the results from a mouse model to human disease pathology is limited by the inherent biological differences, such as lissencephalic brains in mice. Therefore, more refined rodent models and in vitro approaches with cellular models closer to humans are required for the transfer of the results from mouse models to humans.
Since most of the therapies showing promising results in rodents have shown either no or only low efficacy in clinical trials or exhibit severe adverse effects in use, there is currently no established therapy for SSADH-D, and thus no guideline for treatment. From the group of substances directly interfering with the receptor action, only SGS-742, a GABA\(_\alpha\) receptor antagonist, is undergoing a double-blind, cross-over, phase 2 clinical trial with human subjects [73]. Preliminary assessment in patients with mild cognitive impairment showed that administration of SGS-742 improved visual information processing, attention and working memory [71].

The group of classic anti-epileptic drugs, such as vigabatrin and valproic acid, revealed varying outcomes in SSADH-D patients (reviewed in [6]). Vigabatrin, an irreversible inhibitor of GABA transaminase, aims to reduce the high concentrations of GHB by preventing the conversion of GABA to SSA. The drug showed positive effects on behavioral symptoms and refractory epilepsy in only about one third of the patients, and in most cases, the medication has to be discontinued due to loss of vision [6]. Moreover, the outcome in patients is inconsistent, as sporadic cases of worsening of symptoms have been reported [86]. Due to the frequent occurrence of visual field impairment associated with the use of vigabatrin, its use should be thoroughly discussed in such vulnerable patient populations.

### 3.3. Further potential therapy options

#### 3.3.1. Enzyme replacement therapy: special requirements for SSADH-D

Enzyme replacement therapy (ERT) refers to an external substitution of the missing enzyme, usually by delivering a recombinant enzyme by means of injection into the patient’s body. The recombinant enzyme is taken up by the cells and can restore the missing enzyme activity to a varying degree. ERT has been successfully used in various diseases, including several lysosomal storage disorders such as \(\alpha\)-mannosidosis, Pompe disease and classic late infantile neuronal ceroid lipofuscinosis (cLINCL), just to mention a few examples [87, 88, 89].

There are some important, enzyme specific points to consider when developing ERT approaches for a specific disease. The endocytic uptake of the recombinant enzyme is usually dependent on the presence of a suitable receptor protein on the surface of the target cells. In the case of lysosomal enzymes, the delivery is mainly accomplished by the mannose-6-phosphate (M6P) receptors that are ubiquitously expressed. This poses specific requirements for the production of the recombinant enzymes that need to be glycosylated and tagged with M6P to facilitate their uptake into cells. SSADH, on the other hand, is a mitochondrial enzyme whose delivery into its target organelles is coupled with its ribosomal biosynthesis, so that the protein is transferred through the mitochondrial membranes in an unfolded state. Different from lysosomal enzymes, for which uptake mechanism from extracellular fluids exist, SSADH is typically not found outside the cells. Therefore, delivering a recombinant, fully folded SSADH enzyme from extracellular space into the mitochondria of patient tissues poses an extra barrier that needs to be crossed in order to obtain a functional improvement and correct cellular delivery.

Another point to consider is the target organ of the therapy. In lysosomal storage disorders, the brain is frequently affected, and the ERT has to reach the CNS in order to be effective. This raises the question about the route of administration, as it may be necessary to deliver the recombinant enzyme directly into the brain/CNS by means of e.g. intraventricular or intrathecal delivery approaches. This route of delivery is used e.g. in the case of Cerliponase alfa (Brineura\(^\®\)), the recombinant enzyme used in cLINCL [87]. Peripheral administration usually results in poor brain penetration due to the blood-brain barrier (BBB), but may be sufficient in diseases where mainly peripheral organs are affected. Further potential points to consider are the immunogenicity and stability of the recombinant protein used for the ERT. Since the recombinant enzyme needs to be repeatedly administrated into the patient, and a life-long therapy is required, some patients may develop antibodies against the recombinant protein. This has been observed e.g. in the case of Gaucher disease [90]. In some cases, the properties of the recombinant enzyme may be improved by chemical modifications such as PEGylation, as has
been shown for phenylalanine ammonia lyase in phenylketonuria [91], or by packing the recombinant enzyme into carriers such as nanoparticles [92, 93].

In SSADH-D, symptoms of the CNS such as mental retardation and epileptic seizures are profoundly present. However, the exact cause of these symptoms is still under debate. Currently, the main pathogenic role is ascribed to elevated levels of GHB, rather than to GABA. Most of the GHB in our body is produced in the liver, and GHB can efficiently pass the BBB in both directions [94, 95]. Thus, in the case of SSADH-D, it may even be sufficient to clear the periphery from GHB by treating mainly the liver, as the excess brain GHB would be expected to equilibrate with the periphery, thus resulting in clearance of the CNS, too. This hypothesis needs to be tested in the animal model for SSADH-D.

To date, there is only one report on ERT for SSADH-D in the knockout mouse model [70]. Vogel and coauthors produced a glutathione S-transferase (GST)-tagged SSADH fusion protein in a bacterial expression system. After cleavage of the GST tag, 1 mg/kg of the recombinant SSADH was administrated daily in the SSADH knockout mice intra-peritoneally (i.p.), starting on postnatal day 10. The main outcome of the ERT treatment efficacy was the increased survival of the mice. The authors indeed observed a highly significant increase in survival from median of 22 days (PBS-treated mice) to 30 days (rSSADH-treated group) in 4 out of 5 mice in this group. In addition, higher mRNA levels of GABA receptor genes were detected in the treated mice. Highly intriguingly, the authors demonstrated significantly decreased brain GHB levels in the treated mice, implicating that the peripherally administrated recombinant SSADH enzyme was capable of clearing the brain GHB excess [70]. This is an important indication for a possible ERT-based treatment in SSADH-D patients with recombinant SSADH, especially in terms of the route of administration. However, since the recombinant enzyme used evidently does not contain any kind of cell- or brain-delivery sequence, it is unclear if and how the enzyme would be capable of entering the cells, let alone the mitochondria. Therefore, further studies and alternative enzyme delivery approaches should be studied before this therapy can be translated into clinical applications. However, the data of Vogel et al. build an important cornerstone for ERT in SSADH-D, as they show that a reduction of brain GHB concentration may be obtained by peripheral treatment with recombinant SSADH [70].

3.2.2. Gene therapy, stem cell transplantation and genome editing

Already in 2004, Gupta et al. described an adenovirus (AV)-mediated strategy for gene therapy in SSADH-D [96]. Using the SSADH knockout mice, the authors probed two different routes of administration of the virus. Intraperitoneal administration of 10⁸-10¹¹ virus particles by postnatal day 10 resulted in reduction of liver GHB, but had no significant impact on brain or serum GHB concentrations. Retro-orbital injection of 10¹¹ virus particles at day 13 was shown to reduce the GHB concentration in the periphery (liver, kidney, serum) and in the brain, suggesting that this route may have resulted in CNS penetration. Both viral transduction routes increased the liver SSADH activity up to 20% of the wildtype level, and the survival rates of the treated mice were improved, with i.p. administration resulting in better survival rates [96]. However, the observed effects on GHB concentration and enzyme activity appeared to be quite transient, which might be due to the high immunogenicity of AV proteins and consequent elimination of the virus by the host immune system. Despite their capacity to carry large recombinant DNA fragments, adenoviral vectors are not state-of-the-art for human gene therapy anymore, mainly due to their high immunogenicity and highly transient transgene expression in dividing cells. However, the study of Gupta and colleagues has been important in demonstrating that gene therapy is capable of reducing GHB and appears to increase the survival in the SSADH-D animal model [96].

Adenoviral vectors have increasingly been replaced by other viral carriers such as aden-associated viruses (AAV) and lentiviruses. Recent advances in gene therapy and development of a number of gene delivery vehicles based on AAV and lentiviruses have boosted preclinical studies of gene therapy in various genetic diseases, and are currently culminating in numerous clinical trials on gene therapy that have started or are about to start. In fact, Food and Drug Administration (FDA) has recently approved two AAV-based gene therapy vehicles, Luxturna for the treatment of retinitis
pigmentosa and Zolgensma for spinal muscular atrophy (reviewed in [97, 98]). The next couple of years will hopefully result in development of gene therapy approaches for various so far untreatable disorders. For the general developments in AAV-mediated gene therapy, we would like to refer the reader to a recent review by Wang and colleagues [99].

Systemic application of vector particles for gene therapy by intravenous (i.v.) injection may be attractive due to the less invasive route of application. Unfortunately, inefficient penetration through the BBB can compromise gene correction in the CNS. Some viral vectors can cross the BBB, but i.v. injection requires high doses of vector due to the exposition of the vector to circulating antibodies and the widespread transduction of different tissues [100, 101]. Classically, intracranial or intrathecal injection of vector particles is used to circumvent the BBB and has proven to be safe and efficient for the correction of another rare neurotransmitter-related disease, aromatic amino acid decarboxylase deficiency (AADCD; [102]). Here, AAV-mediated gene transfer was used to deliver the DDC gene bilaterally into the putamen of patients.

Potentially, administration of the vectors directly into the cerebrospinal fluid (CSF) may be used to maximize the exposure of the CNS to the vector. It has been shown in non-human primates that this route of application results in a stronger transgene expression throughout the cerebellum and cortex after injection of an AAV9 vector into the cisterna magna compared to i.v. injection [100]. Another problem is the distribution of the transgene at the injection site in the brain. Direct injection into the brain and the limited diffusion rate of viral vectors in the parenchyma lead to only small areas of transduced tissues surrounding the injection site. One solution to overcome this limitation can lie in the use of vectors capable of retrograde transport to distal neuronal projection sites. This can be achieved by using specific envelopes for lentiviral vectors derived from the Rhabdoviridae family or specific AAV serotypes [103, 104, 105, 106].

In addition to gene therapy, stem cell transplantation is considered highly promising for the treatment of many genetic diseases. In many cases, hematopoietic stem cells (HSC) are used for the transplantation. The HSCs can either originate from a suitable donor individual, or patient’s own HSCs can be corrected in vitro (e.g. by gene therapy) and then back-transplanted into the patient in an autologous manner [107]. In the future, it may also be possible to correct the autologous HSCs by genome editing technologies such as CRISPR/Cas [107]. However, due to the unpredictable and sometimes imprecise outcome of genome editing techniques, such as altered splicing or off-target effects [108], these technologies will still require further research before they can be safely applied on patients. Thus, we refrain from further elaboration of these techniques in this review.

The transplanted HSCs contain a reservoir of the functional protein and can deliver the missing enzyme to other cells in the patient organs by means of cross-correction. This mode of action is plausible e.g. for LSDs, in which the missing enzymes can be delivered from one cell to another. Similar cross-correction principle is also utilized by gene therapy (for a review, see [109]). It is thus not necessary to target all cells of the organ, as the cells that have been transduced by the gene delivery vehicle usually produce and secrete an excess of the enzyme that can be taken up by other cells in the vicinity. The degree of cross-correction depends on the fraction of the cells that were successfully corrected, the amount and stability of active, functional enzyme that is secreted, and the uptake efficiency by further cells.

In the case of SSADH-D, both stem cell and gene therapy face similar problems as the ERT approach: mitochondrial enzymes such as SSADH are usually not secreted, and thus cross-correction is unlikely to occur at the level that would substantially correct the deficiency in cells not directly targeted by the therapy vehicle. On the other hand, relatively efficient transduction of the liver can be obtained with certain AAV-based vectors [110, 111, 112, 113]. Thus, if a substantial fraction of the hepatocytes can be corrected by gene therapy, this may result in a large degree of reduction of GHB production, thus lowering the GHB concentration both in the periphery and consequently also in the brain. Since therapy-induced reduction of GHB in the periphery may be a key issue for the development of future therapies for SSADH-D, it would be important to further study the contribution of elevated GHB to the neurotoxic effects in SSADH-D patients.
3.2.3. Small molecules: pharmacological chaperones and read-through drugs

In SSADH-D, the patients exhibit a variety of gene defects including missense and nonsense variants. Although many of these variants are found in a small number of patients, sometimes only in a single family, some defects, such as the substitutions Cys93Phe or Gly409Asp, have been observed in a number of seemingly unrelated families with different ethnic origin. Similarly, some nonsense variants such as Trp204X are present in several patients. Therefore, personalized therapy approaches for specific variants, which are suitable for a number of patients, could be developed for SSADH-D.

Small molecules such as pharmacological chaperones (PCs) for missense variants and read-through therapies for nonsense variants have been developed for other diseases and may also provide novel therapeutic means for SSADH-D. Below, we will briefly summarize these strategies and give examples of their use in other diseases. For a more detailed review on substances used for these purposes, we recommend the reviews by Liguori et al. and Nagel-Wolfrum et al. [114, 115].

In many disorders, the disease-causing variants are not directly located within the active center of the mutated enzyme or within amino acids that are crucial for the protein function, but they rather affect the folding and the tertiary structure of the protein. Such misfolded proteins are frequently not processed and transported correctly, but are detected by the intracellular quality-control systems. This leads to retention of the protein in the endoplasmic reticulum and initiation of the “endoplasmic reticulum-associated protein degradation” (ERAD) pathway, which subjects them to proteasome-mediated degradation [116, 117].

One approach to correct such folding defects and consequently to treat the respective disease is the use of PCs, also known as pharmacoperones [118]. In general, PCs are small molecules that bind to their target proteins and stabilize them. They rescue the function of their targets by facilitating the correct folding, oligomerization and processing, as well as subsequent trafficking of the proteins to their final destination (e.g. lysosomes). In order to be able to do this, the PCs need to bind in a target-selective fashion. PCs are usually not capable of rescuing every single target molecule synthesized in the cells, so that only a fraction of the target proteins will be correctly folded due to binding to the PC. However, this can already lead to a considerable therapeutic benefit, especially in diseases (e.g. lysosomal storage disorders) where a modest increase in the functional protein is enough to produce a therapeutic effect [118, 119].

Examples for PCs that are already applied in humans either as a monotherapy or in combination with e.g. enzyme replacement therapy include migalastat for the treatment of Fabry disease [120, 121], ambroxol for the treatment of Gaucher disease [122, 123], duvoglustat for the treatment of Pompe disease [124] and lumacaftor for the treatment of cystic fibrosis [125]. In addition, Cystadane (betaine) is currently tested in a phase I/II clinical trial (EudraCT Number: 2017-000645-48) for the treatment of aspartylglucosaminuria (AGU) since it has shown PC activity in preclinical experiments [126].

For most of the aforementioned diseases, various kinds of disease-causing variants are known (e.g. missense or nonsense variants, deletions, insertions, splicing defects). However, PCs are mutation-specific and are promising only for some missense variants. Therefore, these substances usually cannot be used for all patients with a given disease, but only for those whose variant is expected to respond to the PC. In all cases, the efficacy of a PC should be tested beforehand in suitable cell culture models, e.g. primary patient cells, for each individual variant. In some diseases, one particular variant is the predominant one found in most patients. In the case of AGU, the most common variant, AGUmiss-major, is responsive to PCs, while other missense variants (e.g. R116W, S72P) do not benefit from the same PC substances [126, 127].

PC therapy holds great promise for the treatment of numerous protein misfolding diseases, but the identification of a suitable PC for a mutated protein is the most challenging part of this treatment approach (reviewed in [128]). In case of misfolded enzymes, most PCs bind to the catalytic site of the enzyme and might even inhibit the activity of the native enzyme. Therefore, in order to avoid inhibitory effects, such PCs should optimally bind in a reversible manner or they need to have a short half-life. In case of receptors (e.g. renal arginine-vasopressin V2 receptor the disease-causing variants of which are associated with Diabetes Insipidus), both agonists and antagonists have been shown to
act as PCs [129, 130]. Furthermore, folding- or oligomerization-defective nicotinic acetylcholine receptors and adrenergic receptors can be rescued by their respective ligands [131, 132, 133, 134]. Other PCs bind to their target proteins in an allosteric way and provide more time for correct folding and processing [119]. In contrast to enzymes and many viral vectors, some PCs are able to cross the BBB, which makes them especially useful for diseases that show a CNS pathology.

In addition to the treatment of conformational disorders caused by mild misfolding, pharmacological chaperones may also be used to enhance the stability and biogenesis of wildtype proteins (reviewed in [118]). This might be especially interesting for diseases that are not caused by folding-defective proteins, but by a shortage of a certain wildtype protein. Due to their beneficial effect on wildtype proteins, the combination of PC and ERT seems promising for some diseases such as Fabry and Pompe diseases [135, 136]. Another application of PCs that targets wildtype proteins is the enhanced production of recombinant proteins, which has successfully been shown for betaine [137, 138].

According to the results of large-scale sequencing projects, 11% of all disease-causing gene defects are nonsense variants that result in an in-frame premature termination codon (PTC) [139]. Hence, special therapies targeting this type of variant, i.e. “nonsense suppression therapies”, have the potential to provide treatment for a substantial number of patients suffering from very different diseases. The first substances known to suppress PTCs were the aminoglycoside antibiotics G418 and gentamicin [140, 141]. For gentamycin, clinical trials have produced variable results [142, 143, 144, 145], as it does not seem to be efficient for all patients. Long-term administration of aminoglycosides is not feasible due to their strong side effects such as ototoxicity and kidney damage [146, 147]. The varying effects of gentamicin may be explained by the fact that gentamicin is not a pure substance, but is composed of different major and minor gentamicin components. In 2017, the minor component gentamicin B1 was postulated to be responsible for the nonsense-suppression effect of gentamicin, but this paper was later on retracted due to problems with the identity of the gentamicin component used in this study [148]. In 2018, the minor gentamicin component X2 was found to be the most active read-through component of gentamicin [149].

In 2007, high-throughput screens for new compounds with nonsense-suppression activity identified PTC124 (Ataluren) as the most promising substance [150]. The luciferase-based high-throughput screening assay that was used for the discovery of PTC124 was later criticized due to the direct effect of PTC124 on firefly luciferase activity [151]. Nevertheless, Ataluren has been used in several clinical trials in cystic fibrosis, Duchenne muscular dystrophy and dystrophinopathy, and it showed some positive effects [152, 153, 154, 155]. The varying efficacy of Ataluren in different diseases may at least in part be explained by the different amounts of residual mRNA that is present in the patient tissues. Depending on the type and location of the PTC, mRNAs harboring nonsense variants may undergo nonsense-mediated decay (NMD), leading to a drastically reduced amount of available mRNA. Hence, NMD inhibition in combination with the PTC suppression will increase the pool of available mRNAs for read-through (reviewed in [156]).

Amlexanox is so far the only substance that combines NMD inhibition and translational read-through [157]. Due to its known anti-inflammatory effects, it has been used for decades in the treatment of other conditions such as ulcerous lesions of the oral mucosa [158]. In cell culture, amlexanox was able to rescue the expression of functional proteins from mRNAs containing nonsense mutations for p53, dystrophin, aspartylglucosaminidase, and type VII collagen [127, 157, 159]. Further potentially efficacious read-through agents include the nucleoside analog clotidine and components of fungi extracts that were identified in a recent drug screen [160, 161]. Since the efficacy of these drugs is also dependent on their capability to cross the BBB, further research will be required to test if these drugs might also provide a therapy option for SSADH-D. Interestingly, the findings of Akaboshi and colleagues suggested that nonsense variants, especially Trp204X and Arg412X substitutions, are frequently present in SSADH-D [31]. Thus, read-through therapies should be tested for nonsense variants resulting in SSADH-D.

4. Future of SSADH-D Research and Role of Patient Organizations
4.1. Current tools for identifying patients and patient registries

As stated above, one major problem in the case of SSADH-D is the difficulty in obtaining the correct diagnosis and identifying as many patients as possible. This will be especially important when novel therapy options hopefully become available in the next years. Fast identification of patients could be obtained by international registries. The International Working Group on Neurotransmitter Related Disorders (iNTD), consisting of 44 partners from 27 different countries, established a web-based registry for inherited neurotransmitter defects, including defects of GABA metabolism [162]. Today, the initiative represents the first longitudinal register for neurotransmitter disorders and enables systematic gathering of data in very high quality. So far, the registry contains data from more than 350 patients. The iNTD collaborates with other networks, such as MetaERN [163], and is part of the Natural History Study of Patients With SSADH Deficiency (ClinicalTrials.gov Identifier: NCT03758521, [164]). The purpose of this multicentric, multinational approach is to collect a large body of clinical data on SSADH-D in order to gain a deeper understanding of the disease progress, symptoms and its molecular consequences. Furthermore, the initiators of this important study also strive at identifying novel biomarkers that can be used in monitoring the disease progress and, importantly, potential therapy success. These data will be essential when considering the approval of future therapy approaches by the national authorities.

In addition to identifying patients through registries and patient organizations, improved diagnosis of new SSADH-D cases is highly important. A large fraction of patients is primarily identified and diagnosed by pediatricians or child neurologists who may not be part of a larger clinical network or a specialized center for rare diseases. Therefore, a highly heterogeneous disease such as SSADH-D might go unnoticed for years. The patients are thus not registered and do not have contact with a patient organization to obtain peer support. Therefore, providing the clinicians with information about SSADH-D as a disease, about the diagnostic means and potential therapies will not only ensure proper care of the patients, but may also result in improved identification of novel cases.

Newborn screening is currently not practiced for SSADH-D. However, as soon as novel (targeted) therapies become available, it would be important to include this disease among those that can be diagnosed as early in the childhood as possible, in order to make sure that the damage caused by the disease is kept at minimum by starting the therapy as early as possible. Very recently, methods addressing the possibility for newborn detection of SSADH-D from dried blood-spots have been developed, paving the way for a screening approach as soon as therapies are available [22].

4.2. Roles of patient advocacy organizations in raising awareness and supporting research

Correct diagnosis is the first key to treating rare diseases. Only when the disease is understood at the molecular level, the clinicians can treat, researchers investigate, and patient advocacy organizations (PAOs) counsel patients regarding further diagnostic investigations and provide support to find appropriate healthcare. Although a particular rare disease may affect only relatively few patients, the number of different rare diseases is currently estimated to be more than 7,000, altogether affecting millions of people. The challenges faced by patients include being correctly diagnosed, informed about possible treatments and receiving proper care and medication. Clinicians, on the other hand, are often confronted with unspecific symptoms, a lack of diagnostic screening instruments and limited access to relevant resources to assist patients with rare diseases. Researchers on the other hand face the problem of obtaining funding and sufficiently large patient cohorts for clinical trials [165]. Furthermore, the availability of validated samples from patients with a specific disease is usually low, limiting the possibilities to use these samples for research work.

General information about a disease is provided through comprehensive databases such as Orphanet [166]. Further professional support comes from large umbrella organizations such as EURORDIS (Rare Diseases Europe) [167] in Europe and NORD (National Organization for Rare Disorders) [168] in the USA. Both organizations play an important role in the drug development process, but further key businesses are advocacy for incentives and patient empowerment. Disease-specific PAOs in turn are significantly smaller, often national patient networks and advocacy groups.
such as SSADH-Defizit e.V. [169] in Germany, De Neu [170] in Spain and the SSADH Association [171] in the US. These disease-specific patient organizations can establish a more direct contact between families, researchers and clinicians. PAOs also provide information on the disease itself and its consequences in the daily life, establishing a platform for exchange with other patients. This is referred to as “peer support” and is the key business of PAOs.

Sadly, rare diseases still receive too little attention, and thus, research projects may be difficult to fund. However, it has been shown that the involvement of patients with rare diseases fosters research. An initiative by EURORDIS only recently published the Rare barometer voices [172]. The survey found that about one third of rare disease patients have taken part in medical research aiming at developing better diagnostic tools and at understanding the underlying disease mechanisms and therapies under development.

Generally, the funding of research on human diseases stems from government agencies (around 30%) and companies (60–65%), with only 5% coming from private resources. In rare disease research, however, non-profit organizations are important sources of funding. For example, in the case of the rare Batten disease, nearly 40% of the funding stems from PAOs, while academic research institutions provide 16% and companies less than 5% of the funding [173]. This is currently also true for SSADH-D. Thus, PAOs for SSADH-D have already started impressively effective seed projects in laboratory-based research as well as clinical trials.

4.3. Future Challenges in SSADH-D

As discussed above, the patient organizations have succeeded to attract further research groups to join the battle against SSADH-D, and a number of novel approaches that directly target SSADH-D at the molecular level are currently under investigation. In SSADH-D, there are no major disease variants that would be present in the majority of patients, but most patients have their own combination of genetic ALDH5A1 variants. Therefore, personalized therapies may have to be developed for single families or even just for a single patient. This requires a detailed characterization of the molecular consequences of the variants in preclinical laboratory studies, followed by a clinical trial with as little as one patient. Although such personalized and variant-specific approaches based on drug repurposing are promising in rare diseases (see e.g. [126, 127]), therapies aiming at benefiting a larger group of patients should also be developed. In the case of SSADH-D, ERT and gene therapy are valid options, despite the limitations due to the nature of the SSADH enzyme (see 3.3.). For the approval of such experimental therapies by the authorities and for the development of suitable biomarkers, the Natural History Study in SSADH-D is of vital importance [164]. The authors are confident that within the next five years, we will see new hope emerging for the treatment of SSADH-D since the researchers are joining their forces to battle this disease.

Author Contributions: All authors participated in writing and editing of the manuscript and agreed with the final version of the article.

Funding: The SSADH-D research in the laboratory of R.T. is funded by SSADH Association and SSADH-Defizit e.V. The work of the Neurotransmitter group at the Centre for Child and Adolescent Medicine in Heidelberg is supported in parts by the Dietmar Hopp Foundation (St. Leon-Rot, Germany), Arbeitsgemeinschaft Pädiatrischer Stoffwechselkrankungen (APS, Germany) and SSADH-Defizit e.V. to T.O.. H.B. has received support from the Physician-Scientist Program at Ruprecht-Karls-University Heidelberg Faculty of Medicine.

Conflicts of Interest: The authors T.O. and R.T. declare to have received funding from SSADH-Defizit e.V, a patient advocacy organization. The funders had no role in the decision to publish the article.
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