Improving the prognosis of nephropathic cystinosis

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Abstract: Cystinosis is an autosomal recessive inherited lysosomal storage disease. It is characterized by generalized proximal tubular dysfunction known as renal Fanconi syndrome and causes end-stage renal disease by the age of about 10 years if left untreated. Extrarenal organs are also affected, including the thyroid gland, gonads, pancreas, liver, muscle, and brain. Treatment consists of administration of cysteamine, resulting in depletion of cystine that is trapped inside the lysosomes. Since cysteamine has a short half-life, it should be administered every 6 hours. Recently, a new delayed-release formulation was marketed, that should be administered every 12 hours. The first studies comparing both cysteamine formulations show comparable results regarding white blood cell cystine depletion (which serves as a measure for cystine accumulation in the body), while a slightly lower daily dose of cysteamine can be used.

Keywords: cystinosis, cysteamine, delayed-release, immediate-release

Introduction

Cystinosis is a lysosomal storage disease. It is inherited in an autosomal recessive manner and is caused by mutations in the CTNS gene, which is located on chromosome 17p13.1 These mutations cause intralysosomal cystine accumulation in cells throughout the body. Today, over 100 pathogenic mutations are known to cause cystinosis. The most frequent mutation found in western Europe is a large deletion of 57,257 base pairs.2

CTNS encodes the lysosomal cystine transporter cystinosin, which is a 367 amino acid lysosomal membrane protein, with seven transmembrane domains. Cystinosin is a proton-cystine cotransporter, coupling the excretion of both cystine and protons out of the lysosome. Protons are carried back into the lysosome by H+-ATPase.3 Besides the cystinosin that is localized on the lysosomal membrane, a second isoform exists due to alternative splicing of CTNS messenger RNA (mRNA). This isoform lacks one of the two lysosomal targeting motifs, consists of the amino acid sequence GYDQL, and was named cystinosin-LKG (the letters LKG representing its last three amino acids). Cystinosin-LKG is localized at the plasma membrane in lysosomes and several other organelles, including the Golgi apparatus and the endoplasmic reticulum. Its exact function remains unclear; however, it is known to facilitate the transport of cystine in an acidic environment, just like its counterpart cystinosin.4

Cystinosis has an incidence of one per 100,000–200,000 live births, and in suspected cases, the diagnosis can be confirmed by measuring white blood cell (WBC) cystine levels and/or by CTNS gene analysis. Three clinical forms of cystinosis can be distinguished, the most frequent being infantile nephropathic cystinosis (OMIM 219800).
Patients generally present before the age of 12 months with polyuria, polydipsia, and failure to thrive caused by generalized proximal tubular damage, also known as renal Fanconi syndrome. It is characterized by increased urinary excretion of several small molecules, including glucose, salts, amino acids, and low molecular weight proteins. Photophobia is caused by corneal cystine crystals, which can be recognized by an experienced ophthalmologist from the age of 12 months. Further, several other organs can be involved during progression of the disease. These include the thyroid gland (hypothyroidism), gonads (hypogonadism, azoospermia), pancreas (exocrine and endocrine insufficiency), muscles (hypotrophy, swallowing difficulties), and the peripheral and central nervous systems. Late-onset or juvenile nephropathic cystinosis (OMIM 219900) is characterized by the same symptoms as infantile cystinosis, but with a disease onset mostly after the first decade of life and a slower rate of disease progression. In contrast, patients with the oculocutaneous form of cystinosis (OMIM 219750) only have photophobia due to corneal cystine crystals while other organs remain spared. The latter two forms of cystinosis are very rare; over 95% of all cystinosis patients are diagnosed with the infantile form.

**Treatment with cysteamine**

Cysteamine (β-mercaptoethylamine) was first introduced as a possible therapeutic agent to treat cystinosis in 1976, and until now has been the only available treatment for the disease. After its introduction as a treatment for patients with cystinosis, cysteamine was first prescribed in the form of cysteamine HCl. Because of the low bioavailability of the hydrochloride salt, it was replaced by phosphocysteamine in 1980. In 1994, cysteamine bitartrate (Cystagon®, Mylan Pharma, Morgantown, WV, USA) was introduced and this is currently the most widely used preparation of cysteamine. Recently, an enteric-coated preparation of cysteamine bitartrate (Procyssbi™, Raptor Pharmaceuticals Inc., Novato, CA, USA) was approved by the US Food and Drug Administration and European Medicines Agency for the treatment of cystinosis.

Cysteamine uses an unknown transporter to enter the lysosome and subsequently breaks the disulfide bond in cystine. This leads to formation of cysteine, which leaves the lysosome using the cystine transporter, and cysteine-cysteamine disulfide, which leaves the lysosome using the PQLC2 transporter (Figure 1). Once in the cytosol, cysteine can be further metabolized to form glutathione, one of the cell’s most potent redox buffers. Indeed, in vitro experiments showed that the altered redox status demonstrated in cystinotic cells was corrected by administration of cysteamine, thereby increasing intracellular glutathione levels.

One hour after ingestion of cysteamine, the drug’s plasma levels reach their maximum while WBC cystine levels are

![Figure 1](https://example.com/figure1.png)  
**Figure 1** Mechanism of lysosomal cystine depletion by cysteamine. Cysteamine enters the lysosome through an unknown transporter and breaks the disulfide bond in cystine. This results in formation of cysteine and a new cysteine-cysteamine mixed disulfide, each of which can exit the lysosome through its own transporter.
at their minimum, followed by a gradual decline in plasma cysteamine levels and a gradual increase of WBC cystine levels. Both reach their original values around 6 hours after ingestion. This underscores the importance of a dosing regimen in which cysteamine is administered every 6 hours; ideally, it should also be administered during the night. Understandably, such a dosing scheme causes a great burden in terms of compliance with cysteamine therapy.

Treatment of cystinosis patients with cysteamine is further hampered by the side effects of the drug. When cysteamine was first prescribed to patients with cystinosis, several of them developed hyperthermia, lethargy, and rash. However, all of these side effects were reversible after the treatment was discontinued, and none of these adverse events are noted today since it has become common practice to introduce cysteamine at a low dose with a subsequent gradual increase over several weeks. Gastrointestinal complaints are due to increased gastric acid secretion after ingestion of cysteamine, and thus concomitant use of proton pump inhibitors is advised. After administration of cysteamine, a small amount is metabolized into the volatile sulfur compounds methanethiol and dimethylsulfide, causing halitosis. Recently, eight cystinosis patients were found to have new adverse events associated with cysteamine, consisting of bruise-like lesions on the elbows, skin striae, and severe bone and muscular pain. Skin biopsies taken from the bruise-like lesions showed reactive angioendotheliomatosis and alterations in collagen fiber diameter. The reactive angioendotheliomatosis could be attributed to increased survival of dermal microvascular endothelial cells after exposure to cysteamine, while the collagen fiber abnormalities could be the result of a synergistic effect of cysteamine and copper deficiency (which was found in all patients identified as having these side effects) on inhibition of collagen cross-linking.

The newly reported side effects of cysteamine resulted in new recommendations regarding dosing. Instead of using body weight to calculate the dose of cysteamine (50 mg/kg/day, with a maximum of 90 mg/kg/day) it is now advised to use body surface area (1.30 g/m²/day, with a maximum of 1.95 g/m²/day). This difference in calculation method is particularly important in growing children, in whom body weight increases faster than body surface area. This may lead to higher (potentially toxic) doses when using body weight to calculate the daily cysteamine dose, since 90 mg/kg may exceed 1.95 g/m².

Despite its side effects and four times daily dosing regimen, cysteamine remains the cornerstone of treatment for cystinosis and adequate administration of cysteamine significantly reduces the rate of progression towards end-stage renal disease, and postpones or even prevents the occurrence of extrarenal complications. Hypothyroidism, myopathy, pulmonary dysfunction, and diabetes mellitus (caused by pancreas insufficiency) occur less frequently in cystinosis patients who have been treated with cysteamine for a prolonged period of time. Further, while children with cystinosis show more pronounced growth retardation than children with other renal diseases, adequate treatment with cysteamine significantly improves growth parameters. Nonetheless, most children with cystinosis also require treatment with recombinant human growth hormone to achieve a final height higher than two standard deviations below the mean.

Unfortunately, not all complications of cystinosis are delayed by administration of cysteamine. Cysteamine does not influence the incidence of primary hypogonadism in male patients with cystinosis, nor does it improve delayed puberty. Also, male infertility due to azoospermia is not ameliorated by adequate treatment with cysteamine. Further, systemic administration of cysteamine does not prevent formation of corneal cystine crystals, indicating a need for topical cysteamine eye drops which should be applied several times daily. Recently, a new gel formulation of cysteamine (Cystadrops®, Orphan Europe, Montluçon, France) was tested for its ability to diminish the formation of corneal cystine crystals, with better effects and less applications needed per day compared with the usual cysteamine hydrochloride formulation.

### Other novel treatment strategies

The most promising results with regard to treatment of cystinosis without use of cysteamine come from the field of stem cell transplantation. In a CTNS−/− mouse model treated with wild-type allogeneic bone marrow and hematopoietic stem cells, the mice that received a bone marrow graft showed better renal function after 4 and 15 months, as well as decreased cystine content in a number of tissues, including the brain, eyes, heart, kidneys, liver, skeletal muscle, and spleen. However, despite these promising results, tissue cystine levels remained higher than those in asymptomatic heterozygous carriers, so administration of cysteamine may still be warranted.

In addition to bone marrow transplantation, gene therapy was also studied in the CTNS−/− mouse model in order to minimize the use of immunosuppressive agents, along with the risks of rejection and development of graft versus host disease. Hematopoietic stem cells from CTNS−/− mice were
transplanted with the human CTNS complementary DNA (cDNA) sequence followed by transplantation into CTNS\textsuperscript{-/-} mice. Again, tissue cystine levels were significantly decreased after 4 and 8 months, including in the brain, eyes, kidneys, liver, muscle, and spleen, but were still higher than those after wild-type bone marrow transplantation.\textsuperscript{37}

Prodrugs are under development in order to diminish the bad taste and smell of cysteamine and to increase its half-life. Several prodrugs of cysteamine have been tested in vitro;\textsuperscript{38,39} however, none are as effective as cysteamine in decreasing intracellular cystine levels, and none have been tested in vivo.

**Delayed-release cysteamine bitartrate**

The observation that administration of cysteamine directly in the small intestine resulted in higher plasma cysteamine levels with a higher area under the curve when compared with gastric administration of the drug\textsuperscript{40} led to development of the enteric-coated formulation of cysteamine bitartrate (Procysbi). In contrast with the immediate-release formulation of cysteamine bitartrate, this new delayed-release formulation can be administered every 12 hours instead of every 6 hours.\textsuperscript{41} The delayed-release formulation consists of capsules containing small granules than can be administered through a feeding tube. This is important, given that most small children with cystinosis require a lot of fluids and electrolyte supplements, and thus have feeding tubes that are also used for administration of medication.

In a Phase III trial testing the noninferiority of the delayed-release formulation, 41 nontransplanted patients with cystinosis were randomized to treatment with either immediate-release or delayed-release cysteamine bitartrate for 3 weeks and then switched to the other formulation for a further 3 weeks. The mean difference in WBC cystine levels after treatment with the delayed-release formulation compared with the immediate-release formulation was 0.08 nmol 1/2 cystine/mg protein (95.8% confidence interval 0.00–0.16), meaning that the primary endpoint of noninferiority was met.\textsuperscript{42}

Interestingly, the average total daily steady-state dose of the delayed-release formulation was 82% of the previously established incoming dose of the immediate-release formulation,\textsuperscript{42} indicating that similar results for depletion of WBC cystine can be achieved with a slightly lower daily amount of cysteamine. In theory, this could result in fewer side effects. An early pilot study was conducted comparing the excretion of volatile sulfur compounds in four patients after ingestion of immediate-release or delayed-release cysteamine bitartrate (as part of the crossover Phase III clinical trial). This study showed a trend towards a lower area under the curve for dimethylsulfide excretion in all four tested patients after ingestion of the delayed-release formulation. However, this trend did not reach statistical significance, patient numbers were small, and the measured levels of dimethylsulfide remained above those that cause halitosis at all time points.\textsuperscript{43}

A long-term study is presently underway using delayed-release cysteamine bitartrate in cystinosis patients with renal Fanconi syndrome and in those who have undergone renal transplantation. Again, as part of this study, excretion of volatile sulfur compounds in breath will be measured in a subgroup of patients, comparing the levels at study onset when patients are treated with the immediate-release formulation and again a few months after treatment with the delayed-release formulation. The final results of this study are pending, but preliminary data show that dimethylsulfide breath levels again remained persistently above the threshold that causes halitosis; however, further analysis should determine whether the area under the curve is influenced by the delayed-release formulation. Previous reports regarding the long-term treatment of two cystinosis patients with the delayed-release formulation show a good effect on the mean trough WBC cystine level, with no deterioration of renal, thyroid, or liver function observed during a treatment period of 6 years.\textsuperscript{44}

**Prognosis**

Overall, the quality of life for patients with cystinosis has improved significantly since the early 1980s. Since the use of cysteamine to treat cystinosis and the introduction of renal replacement therapy in pediatric populations (either through dialysis or by renal transplantation), patients with cystinosis are increasingly surviving into adulthood, whereas before the introduction of cysteamine as a possible treatment for cystinosis they generally died from end-stage renal disease by around the age of 10 years.\textsuperscript{27} This increased life expectancy has caused an increase in the number of longer-term complications of the disease, that were not noticed in younger patients. This increased interest and recognition of longer-term complications of cystinosis prompted a search for a possible treatment for these complications. For instance, male infertility due to azoospermia was first described only a few years ago, awareness of which has enabled patients to seek further treatment options, such as testicular sperm extraction, which could open the possibility for male patients to produce their own offspring.\textsuperscript{8} Another example is the severe growth retardation seen in patients with cystinosis. Since this is now
treated more effectively (eg, by improving nutritional and electrolyte status, prescribing cysteamine, and administration of recombinant human growth hormone), today most cystinosis patients can reach a final height within the normal range, which further improves their quality of life.

Apart from better recognition and treatment of (late-term) complications of cystinosis, better treatment of the disease also improves the long-term outcome for the patient. Cystinosis is now more frequently recognized at a young age, resulting in more patients being adequately treated with cysteamine at a young age. This is very important, since early initiation of cysteamine therapy can significantly postpone or even prevent complications of the disease, including deterioration of renal function, hypothyroidism, and myopathy. However, one of the most important burdens regarding cysteamine therapy is the need to administer the drug every 6 hours in order to ensure adequate WBC cystine depletion. Development of the new delayed-release formulation can overcome this problem, which will likely improve patient compliance with therapy, resulting in fewer long-term complications of cystinosis and improving patient life expectancy even further.

It remains uncertain to what degree potential new therapies for cystinosis (including bone marrow transplantation and/or gene therapy) will further improve the long-term outcomes. Although impressive results have been documented in the CTNS−/− mouse model, there are as yet no results of such treatment in humans.

Conclusion

From 1976 until now, cysteamine has been the only treatment available for cystinosis. The advent of cysteamine has significantly improved the outcome for patients with cystinosis, as have the better recognition and treatment of the complications of the disease. New treatment options, such as bone marrow transplantation are promising, but are not currently performed routinely in humans. Further, although intracellular cystine levels in a number of tissues in CTNS−/− mice decrease significantly after bone marrow transplantation derived from their wild-type litter mates, cystine levels remain above those of asymptomatic heterozygous carriers.

Thus, even after bone marrow transplantation, there may still be a need for low doses of cysteamine to decrease the remaining accumulation of cystine. This indicates a need to further improve the treatment of cystinosis with cysteamine. In this regard, the new delayed-release formulation of cysteamine bitartrate is of importance to improve compliance with cysteamine therapy.

Disclosure

The authors report no conflicts of interest in this work.

References


