Cystinosis: Therapy adherence and metabolic monitoring in patients treated with immediate-release cysteamine

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ARTICLE INFO

Keywords:
Adherence
Cysteamine
Cystine level
Cystinosis
Metabolic monitoring

ABSTRACT

Background: Cystinosis is a metabolic disease caused by intracellular accumulation of cystine within lysosomes. Development of symptoms can be delayed significantly by a life-long therapy with cysteamine, a drug that enters the lysosome and reacts with cystine thereby enabling its export from the organelle.

Method: During a period of 16 years, blood samples of 330 cystinosis patients were analyzed to investigate therapeutic adherence and metabolic control in patients treated with immediate-release cysteamine. The accepted therapeutic goal is to measure intracellular cystine levels in white blood cells every 3 months and to keep them below 0.5 nmol cystine/mg protein (= 1 nmol hemicystine/mg protein).

Results: 42% of measurements were within the desired 3-month interval, 38% were done every 3–5 months, 11% every 6–8 months, 5% every 9–12 months and 4% after a 12-month interval only. 64.4% of the measurements were higher than the therapeutic target value. Median cystine levels increased with longer control intervals.

Conclusions: The majority of the cystinosis patients showed insufficient metabolic adjustment. Intraocular cystine levels were not done as often as recommended and were not within therapeutic range. Poor therapy adherence is likely to be caused by gastrointestinal side effects of immediate-release cysteamine. Incorrect intervals between drug intake and blood sampling could contribute to the results.

1. Introduction

Cystinosis is an autosomal recessive metabolic disease characterized by reduced cystine export from lysosomes due to absent or abnormal cystinosin, the integral lysosomal membrane cystine transporter [1]. This results in storage of lysosomal cystine, which is rapidly absorbed from the gut and reacts inside the lysosomes by intralysosomal protein degradation and rapidly forms the mixed disulfide cysteine-cysteamine. This product can be exported out of the lysosomes by the cationic amino acid transporter PQLC2, thereby decreasing the cellular cystine level [6].

Cystagon® (Mylan Pharmaceuticals, Canonsburg, PA) is an immediate-release form of cysteamine and contains 50 or 150 mg of cysteamine per capsule as mercaptamine bitartrate. It received the FDA approval in the USA based on clinical data after various clinical tests in 1994 and is taken orally 4 times per day in regular intervals of 6 h immediately after or together with a meal. After 2 h, the cellular cystine level drops to a minimum, and rises again to its original high level after 6 h [7]. Until the age of 12 years, standard daily dose is 1.30 g/m² body surface area. In older patients over 50 kg, the recommended daily dose is 2 g up to a maximum of 1.95 g/m². Monitoring of the cystine concentration in leukocytes at specific intervals is required for correct adjustment of the dose. The drug must be taken life-long. The therapeutic target level in white blood cells (WBCs) is ≤0.5 nmol cystine/mg protein (<1.0 nmol hemicystine). Cysteamine therapy can delay disease progression very efficiently [8].

Many patients experience considerable side effects upon cysteamine treatment: mouth and body odour occurs as well as gastrointestinal problems like frequent nausea and vomiting, abdominal discomfort and...
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0.5 nmol cystine/mg protein. While the results of 3-month follow-ups
intervals in the total group. The dotted line indicates the target value of
classical method by Spackman, Stein and Moore [13], a method with
2.1. Cystine measurements
The cystine measurement of the leucocytes was carried out by the
classical method by Spackman, Stein and Moore [13], a method with
ion chromatographical separation of amino acids by a cation exchange
column and a step gradient. N-Ethylmaleimide (NEM) was added to
capture free sulphydryl groups. Two internal standards were measured in
addition (norvaline and cystine).

3. Results
A total of 42% of the controls were within the desired 3-month interval,
38% were done every 3 to 5 months, 11% every 6 to 8 months,
5% every 9 to 12 months, and 4% were done on after more than
12 months (Fig. 1).

Fig. 2 shows intracellular cystine levels within the respective time
intervals in the total group. The dotted line indicates the target value of
0.5 nmol cystine/mg protein. While the results of 3-month follow-ups
and follow-ups from 3 to 5 months show barely any differences (the
median was 0.6 nmol cystine/mg protein each), an increase in cystine
levels was observed with longer intervals (e.g. > 12 month with a
median of 0.8 nmol cystine/mg protein). Longer monitoring intervals
go along with higher mean intracellular cystine levels. The many out-
liers found in the first two interval groups are remarkable, especially
since some of them are equal to untreated cystinosis.

However it must be considered that longer intervals between the
measurements may result in less frequent adjustment of the medication
dose. Even patients who were following the strict medication schedule
could suffer from a underdosing represented by the highest median of
their higher interval group.

The intra-individual fluctuations of the 10 patients with the most
frequent examinations during the mentioned period are illustrated in
Fig. 3 with 63 blood measurements the first patient and 45 samples in
the last patient. The medians were between 0.40 and 0.79 nmol cy-
stine/mg protein. The best result could be found in patient 1 with 68%
of blood samples below 0.5 nmol cystine/mg protein. Despite of two
outliers (max. 1.22 nmol cystine/mg protein), 50% of the other 61
blood samples were located between 0.3 and 0.53 nmol cystine/mg
protein. Patient 2 had many mild and extreme outliers. Levels of up to
3.6 nmol cystine/mg protein were achieved at times. Patient 4 (47
blood samples) is represented by the 4th box which is the widest one
and shows a high degree of fluctuation and the highest median (almost
0.8 nmol cystine/mg protein). Recommended cystine levels could not
be achieved permanently in any of the patients.

The median of all measurements was 0.63 nmol cystine/mg protein,
and therefore above the therapeutic target value (Fig. 4). Including the
middle 50% of the data, the box is characterized by an interquartile
range of 0.43 till 0.9 nmol cystine/mg protein. The lowest cystine level
was 0.04 nmol cystine/mg protein, the highest could be found at
8.9 nmol cystine/mg protein. All together 64.4% of the measurements
were over the target value of 0.5 nmol cystine/mg protein, only 35.6%
reached the desired range. It shows the extreme outliers, which were
found both in patients who seemed therapy-adherent as well as in pa-
tients who were less compliant with follow-up measurements.

Fig. 5 demonstrates the relative distribution of cystine levels mea-
sured. It shows that 35.6% of the levels were below 0.5 nmol cystine/
mg protein, which means that the metabolic monitoring was in-
sufficient in 64.4% of the samples. 28.2% of the levels were > 0.5
and ≤ 0.75, 16.8% > 0.75 and ≤ 1, 6.7% > 1 and ≤ 1.25. The last
three categories show barely any differences: levels > 1.25 and ≤ 1.5
could be found in 4.1% and levels > 1.5 and ≤ 2 and > 2 were
reached by 4.3% each.

Fig. 6 presents the subdivision of the measurements in the different

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**Fig. 1.** Time intervals between two blood samplings of cystinosis patients for
WBC cystine content measurements (3278 blood samples of 162 patients). The
appointments were considered individually and independent of the corre-
sponding patient.

**Fig. 2.** WBC cystine levels measured in the respective time intervals, in-
dependent of individual patients to illustrate a correlation between appoint-
ment adherence and cystine level (All together: 3349 blood samples). The
dotted line indicates the target value of 0.5 nmol cysteine/mg protein. The grey
coloured boxes represent the middle 50% of the data, the band inside the boxes
shows the median. Stars represent the extreme outliers and circles the milder
ones.
age groups. Values obtained within the first 12 months of age were assigned to the category Newborn/Infant, from the start of the 2nd year to the end of the child’s 3rd year, the category assigned is Small Child, from the start of the child’s 4th year until the end of the 12th year, the category is Child, from the start of the 13th year to the end of the 18 year – Youth, and thereafter the category Adult. It can be seen that the median values of Small Children, Children and Youths were almost identical (0.61 to 0.60 to 0.62 nmol cystine/mg protein). The median value was highest for infants at 0.95 nmol cystine/mg protein, followed by the value for adults at 0.7 nmol cystine/mg protein. No statistical outliers are found in the newborn category; the total number of included values is, however, at \( n = 24 \) the lowest, whereas the children group accounts for most measurement data (\( n = 1374 \)). The adult category showed the highest outliers, and the interquartile range was also larger, similarly for the newborn group.

4. Discussion

White blood cell cystine levels have to be kept all-day-long below a threshold level in order avoid cystine crystal formation and subsequent cellular damage. Several studies monitoring cystinosis patients receiving cysteamine therapy showed that adherence is often insufficient to achieve a stable cystine level below the recommended therapeutic target value. This may be due to the strict medication schedule requiring getting up at night to adhere the proper 6 h dosing interval and to the side effects of the medication, such as bad smell and gastrointestinal problems [7,9,10]. Approximately 14% of the affected patients do not tolerate cysteamine therapy due to strong nausea and vomiting, which worsens compliance [5]. If the drug is tolerated, only 23% of patients comply with the strict medication schedule, 17% medicate only during the day. However, the cystine concentration will greatly increase if the medication is taken with a small delay of 2–3 h [7].

The current study demonstrates that metabolic control in a large cohort of cystinosis patients taking immediate-release cysteamine seems to be poor. Intra-individual fluctuations in different blood samples were high and were not rarely above 1 nmol cystine/mg protein in WBCs. Fluctuations may be due to the deviation from prescribed intake times of 6 h prior to drawing blood or an inaccurate or missed dosage. Due to the pharmacokinetics of immediate-release cysteamine, strict adherence of the blood sampling to the 6-h interval is mandatory.

As an alternative to Cystagon©, which is released from the capsule in the stomach, a drug called Procyshin© (Raptor Pharmaceutical Corporation, USA) was launched in the market, which is released and
absorbed in the small intestine due to its enteric coating and shows improved cysteamine pharmacokinetics. It allows reducing the dose required with Cystagon© to 82% and only needs to be taken twice daily [10]. The simplified intake of that drug and fewer gastrointestinal side effects could improve adherence.

Cystinosis patients require life-long care and monitoring of their intracellular cystine levels, which generally requires time and the willingness of having blood drawn. Therefore, it is not surprising that patients, primarily in those examined for many years, in part over the entire period of 16 years, get tired of keeping follow-up appointments. But results show also that strong fluctuations in WBC cystine levels occur even in dedicated patients, who are under regular care.

No information was provided about the prescribed dosage with each patient, the actual dose interval or the time the medication was administered last before the blood sample was taken. If the predefined interval was exceeded the result would be a rise in cystine measurements. Another way how cystine levels could be falsely low would be if the requested 6-h interval between the last cysteamine administration and the blood sampling was shorter.

Whereas this retrospective analysis cannot correlate metabolic control with clinical outcome, it clearly shows insufficient metabolic control or wrong intervals between drug intake and blood sampling with immediate-release cysteamine treatment. In order to delay disease progression ever further, greater care has to be taken with respect to adequate dosing and correct blood sampling.

Declaration of Competing Interest

None declared.

Acknowledgements

T.M. received consulting fees and research grants from Orphan Europe (France; No. 141217) and from Horizon Pharmaceuticals (Ireland; No. 20150904) and acknowledges support from the Open Access Publication Fund of the University of Muenster (Germany).

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Fig. 6. Measurements in the different age groups. The age of the patient is determined at the follow-up appointment and categorized accordingly.