PAUL AND POMPE DISEASE

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AND POMPE DISEASE

Pompe disease is a rare genetic disorder which can cause a range of health issues. The most common symptom is muscle weakness which deteriorates with time if the disease is left untreated.

This brochure explains in simple terms the cause of Pompe disease and which symptoms may arise at a young age.

This document has been organised and funded by Sanofi and is intended for children diagnosed with Pompe disease.

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Hi, my name is PAUL and I am 8 years old!

I live on a farm with my parents and little brother, P[T[R]].

WE HAVE LOTS OF COWS, CHICKENS AND OTHER ANIMALS

During the summer holidays, I like to help my mum and dad with taking care of the animals.

I sometimes find it hard to carry heavy things because my arms DON'T FEEL STRONG ENOUGH.

MY MUM SAYS THAT'S BECAUSE I'M STILL VERY YOUNG



Just two more weeks and I'll be going back to $\$

I'm going to be in year 4 this year which means that I will be one of the big kids at school now. But for now, I get to play in the HAYLOFT with my little brother, Peter.

Except I have trouble climbing the ladder

BECAUSE IT'S SO TIRING



My legs also feel so TIRED



I feel so weak and tire so quickly that my parents are starting to worry about me. So we went to visit the doctor in the village.

We spoke to the doctor and he did a couple of tests.





He said that I was probably run down from helping at the farm all day.

I just needed to slow down a little and get some rest and I would be fine.



School's started again but I'm too tired to go to school all day.



I HAVE TO STAY AT HOME NOW

My parents don't believe that I'm just worn out and my mum went and looked on the internet to see what else it could be.

We called the doctor again and we made an appointment with the paediatrician – that's a long word for a doctor who treats children. When we saw the doctor at the hospital, I told her all the things I had been having trouble with:







She said that she thought she knew what the problem was.

THE DOCTOR CALLED US AS SOON AS SHE HAD THE RESULTS



We went back to the hospital for the results.

The doctor told us that I had MPE SEASE and explained to us what this disease is.





My body is just like my room:

WHEN IT GETS MESSY, IT NEEDS TO BE CLEANED

The problem is that my body doesn't have enough '([[AN[R]'] so the 'M[S]' piles up and causes all the problems I've been having.

Some people even have so few '(LEANERS' that they have trouble with this disease when they're only a baby.

The doctor said to me:

"You will need to come to the hospital regularly for check ups. By visiting us, it means we can help you to MANAGE THE DISEASE."

There are other children with this disease who are also being treated in this hospital. When we got back home, my parents were relieved that the doctors knew what was wrong and were ABLE TO SUPPORT ME

