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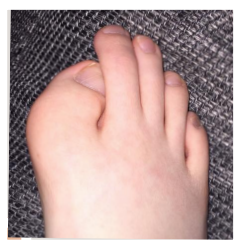
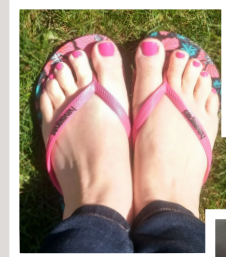
FOP Awareness Day 2022

23rd April

WHAT WILL YOUR FEET WEAR?

Fibrodysplasia ossificans progressiva, or FOP, is one of the rarest and most disabling conditions known to medicine: there is no known treatment or cure. Over time, bridges of extra bone develop, forming a second skeleton that imprisons the body: a healthy mind locked inside a frozen body. It can cruelly take a child's ability to walk, run or play overnight. There is no way to stop its progression.

Help us to help those affected by this devastating disease.



Share your wackiest or most stylish toes!
#FunFeet4FOP



FOP Friends is a registered charity: England and Wales 1147704 and Scotland SC046950





Finding a cure for FOP

What?

When?

Where?

For more information...



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Finding a cure for FOP

You smashed it!

Thanks to your support, the

raised an amazing...

£

Thank you for
being a Friend!



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