October 21, 2019

Welcome from the Chair:

The WMFC has been helping Waldenstrom's Macroglobulinemia patients and their families in Canada for the last 20 years and has made a huge difference to patients and their families as they take on their unique WM journey.

In the late 1970's my mother developed WM and had no treatments other than blood transfusions. She passed away about six years later. This is in remarkable contrast to today. I was diagnosed in 2016, have had treatment and am looking forward to another 20 years of good health.

The WMFC is here to help all WM'er's in Canada. We all need information and support of others to manager our own journeys. Please reach out to any Board member, and we will do all we can to assist you and your family. Whatever it is you need, we will try and find it for you.

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Imagine a Cure! Waldenstrom's Macroglobulinemia Foundation of Canada
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