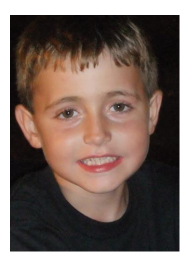


Don't Let Duchenne Derail Another Boy's Life~Stop Duchenne In Its Tracks

Dear Friends of John Owen's Adventure!

We humbly ask for your support and be part of our <u>12th Picnic in the Park</u> with your donation.



Enclosed is our 2019 pledge request, we ask that you to continue to be part of history as we Derail Duchenne Muscular Dystrophy. We are in need of **products, gift cards and basket items** to support our event that attracts **2,000 guests on July 13, 2019**. Any donation big or small is **greatly appreciated**.

JOA was started in 2008, after our son, John Owen Dumm, was diagnosed with Duchenne, much progress has been made since his diagnosis, including the first approved treatment of Duchenne, Exondys 51. Not a cure, but certainly a step forward, however, <u>much more work needs to be done to stop this muscle killer</u>.

That is where you come in!

JOA is a 501c3 and your donation is 100% tax deductible *EIN 26-2672169*. JOA's mission is to end this most common, fatal muscle disease that affects 1 in 3,500 boys.

Thank you for taking the time to read this letter and for your <u>heartfelt consideration</u> in helping us Derail Duchenne! Please feel free to reach me at 440-570-2167.

We will be honored to accept your donation at any time.

Best regards,

Jen Dumm

We are so very grateful to our constant corporate sponsors and their Awesome Generous Spirit to Derail Duchenne! Join Them, Will You?

There are many other amazing corporations joining our fight, we could not fit them all THANK YOU!



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