The Patient Voice

Ryan Fischer
SVP Community Engagement
“The value placed on the benefits and risks of any treatment can only truly be understood by the patients/caregivers themselves, for it is they who have to make the treatment choices and it is they who bear any associated risks.”

(~ quote from a health economist at a recent conference)
Gap between scientific data and patient experience

Your Voice: Bridging the gap

- Disease burden & unmet need
- Outcome measures and endpoints
- Preferences w/ risks, benefits, uncertainty
- Symptom priorities
- Quality of life
- Meaningful benefit
Patient/Caregiver Data Takes Many Forms…

**Qualitative Data *(Describes)*
- Focus Groups
- Open Ended Survey Questions
- Testimonies
- Interviews (in person or phone)
- Town halls

**Quantitative Data *(Defines)*
- Patient Registries
- Patient Reported Outcomes
- Quality of Life Instruments
- Polling data
- Surveys

**Preference Data** *(can do both)*
Preference Studies – Quantifying the patient voice

5 studies completed to date
• Caregivers preferences
• Symptom priorities
• Meaningful pulmonary outcomes
• Multi-stakeholder preferences
• Preferences for emerging gene therapies

Underway
• Global preferences for treatments

Academic collaborators
Dr. John Bridges (The Ohio State)
Nonie Crossnohere (JHU)
Holly Peay (RTI International)

Funding and participation on advisory groups:
• Pfizer
• Everylife Foundation
• Santhera Pharmaceuticals
• Solid Biosciences

#PPMDConference
### Stated Preference Methods

- Methods for collecting and analyzing data about what people think and feel.

  Quantitative preference surveys: provide numbers to explain
  - How much 'bad' will people accept for the 'good'?
  - How much risk will a person tolerate given a particular benefit?
  - How do people prioritize among important factors when making a decision?

<table>
<thead>
<tr>
<th>Task 2</th>
<th>Drug A</th>
<th>Drug B</th>
</tr>
</thead>
<tbody>
<tr>
<td>Does it slow disease progression?</td>
<td>Slows for 5 years</td>
<td>Slows for 3 years</td>
</tr>
<tr>
<td>How many people would benefit?</td>
<td>75%</td>
<td>25%</td>
</tr>
<tr>
<td>What is the extra risk of kidney damage?</td>
<td>No extra risk</td>
<td>20% extra risk</td>
</tr>
<tr>
<td>What is the extra risk of fracture?</td>
<td>20% extra risk</td>
<td>No extra risk</td>
</tr>
<tr>
<td>Which drug would you choose?</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Consistent themes learned from the data

• Stopping or slowing disease progression are *both* valued treatment outcomes, patients and caregivers are willing to take on risk and uncertainty to achieve these outcomes.

• The community prioritizes protecting muscle function and treatments targeting secondary symptoms of cardiac and pulmonary function most and are willing to take on risks, burdens, and uncertainties for benefit.

• An appropriate balance of benefits and risks may be different for people with Duchenne or Becker, their caregivers, and the professionals who develop treatments and manage medical care.

• Decision making for clinical trial participation is mainly driven by the chance for benefit (skeletal, cardiac, pulmonary) over risks, burdens and uncertainties.
Why this matters

• These studies allow us to engage a large group of community members in patient centered research

• Your contributions to these studies enable us to better communicate your preferences to a range of stakeholders in the drug development ecosystem

• Preference change over time, we will continue doing these studies in order to accurately reflect the current environment

  • Global study underway (6 countries)
  • Thank you to everyone who has contributed data!
THE DUCHENNE REGISTRY

THE STRENGTH OF THE REGISTRY IS YOU

Your data is critical in the fight to end Duchenne

Parent Project Muscular Dystrophy
The Duchenne Registry

- Online self-reported registry for individuals with Duchenne or Becker, as well as carrier females

- Established by PPMD in 2007, in collaboration with Emory Genetics and the Centers for Disease Control

- Largest patient reported registry globally

#PPMDConference
Inform understanding on NH of DBMD
- Family Hx and Location
- Genotype/phenotype
- Care interventions
- Collect PROs on disease

Patient Resources
- Free genetic counseling
- Free genetic testing
- Clinic trial education
- Trial Finder
- Decision making tool

Drug Development
- Recruit for studies
- Prep to trial
- PROs on standard of care

Share information
- Individuals with DBMD
- Families and caregivers
- Healthcare providers
- Researchers
- Industry
- Regulators
- Payers

REGISTRY AIMS AND GOALS
5000+ Registrants

Diagnosis

- Manifesting carrier (I am a carrier of Duchenne or Becker and I have symptoms)
- Possible carrier (I do not know for sure if I am a carrier)
- Confirmed carrier (I am a carrier of Duchenne or Becker and I do not have any symptoms)
- Duchenne or Becker (not clear yet)
- Becker
- Duchenne

76%

Mutations

<table>
<thead>
<tr>
<th>Mutation</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deletion</td>
<td>64.6</td>
</tr>
<tr>
<td>Nonsense</td>
<td>12.5</td>
</tr>
<tr>
<td>Duplication</td>
<td>9.6</td>
</tr>
<tr>
<td>Deletion – small mutation</td>
<td>5.4</td>
</tr>
<tr>
<td>Splice site</td>
<td>3.9</td>
</tr>
<tr>
<td>Duplication small mutation</td>
<td>2.2</td>
</tr>
<tr>
<td>Insertion</td>
<td>0.8</td>
</tr>
<tr>
<td>Missense</td>
<td>0.5</td>
</tr>
<tr>
<td>other</td>
<td>0.4</td>
</tr>
</tbody>
</table>

Registry Modules

<table>
<thead>
<tr>
<th>Genetic information</th>
<th>Behavior</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medications</td>
<td>Bone Health</td>
</tr>
<tr>
<td>Muscle Function</td>
<td>Family History</td>
</tr>
<tr>
<td>Insurance</td>
<td>Pain</td>
</tr>
<tr>
<td>Cardiac</td>
<td>Steroids</td>
</tr>
<tr>
<td>Respiratory</td>
<td>Diagnosis</td>
</tr>
</tbody>
</table>
10-Year Registry Report

IMPACT

- Over 5000 Registrants
- 115 countries
- 100+ Trials and Studies Recruited
- Data collected over 12 years

#PPMDConference
The NEW
Duchenne Registry
Platform
(app based)
Set to launch late summer

In partnership with
THREAD™
Welcome

This walkthrough will explain the goals of The Duchenne Registry, the impact it may have on your life and will allow you to provide your consent to participate.

Learn more about how the consent process works.

Learn More

To start an activity, select from the list below:

- Diagnosis Module
- Muscle Function
- Corticosteroid Module
- Breathing/Respiratory Module
- Bone/Orthopedic Module

Activity Complete!
Consent Process

- Quick and straightforward consent process to allow for your de-identified (anonymous) data to be used for research purposes
The Interchange combines patient-reported data, clinician-reported data (on approved therapies), and data from electronic health records (care visits), to analyze and understand real world evidence about the Duchenne and Becker progression.

- PPMD partnered with Sarepta Therapeutics to launch the Interchange
- Data from the Duchenne Registry is the first patient reported data included in the Interchange
- We are looking to marry the PRO data with Electronic Health Records.
- Data from patients on Exondys 51 (eteplirsen) will be the first post-market surveillance data in the Interchange
- The Interchange is built as a model for additional Industry partners to be added on after the pilot of Exondys 51 (Sarepta’s EVOLVE Registry)

#PPMDCconference
PPMD approach to capturing Electronic Health Records with Clinical Care Centers –

How PPMD will enhance the new Duchenne Registry

You will complete an informed consent in the new PPMD THREAD platform giving PPMD permission to pull Electronic Health Records

Consented List
Patient A
Patient B
Patient C

PPMD prepares a list for each institution of all consented patients seen at that institution

Institution shares with PPMD data elements in EHR for these patients through a secure API

HIPPA Compliant Data Transfer

Data is ingested into new Interchange

Duchenne Interchange
Example of how it works:

Patient consents with The Duchenne Registry and “Springfield Hospital” to push their EHR data to Interchange.

(EHR data elements Patients from “Springfield Hospital”)

Doctors at “Springfield Hospital’s” are provided access to secure Data in Interchange (PPMD PRO + EHR data elements)

The good news for clinics! We are NOT asking clinics to enter anything into a new database.
• The Evolve Registry is owned and operated by Sarepta and is an important and exciting project to gather real world evidence data in DMD patients who are on their approved medications.

• The Evolve Registry has started enrolling sites and recruiting patients.
Duchenne Interchange

- THE DUCHENNE REGISTRY
  - Patient Generated Data

- Electronic Health Record (EHR) Data

- Clinician Generated Data

- PPMD Duchenne Registry PRO Data

- Sarepta EVOLVE DATA

- EHR DATA FROM CLINICS

POST APPROVAL DATA

DUCHENNE OUTCOMES RESEARCH INTERCHANGE

Analytics

#PPMDConference
What do I need to know and do?

- The NEW Duchenne Registry App is launching this fall!
  
  - Visit The Duchenne Registry at Interaction Alley at Conference!
  
  - Be sure to indicate with us which clinic you are currently receiving care
  
  - Provide us with the most up to date email for your account(s)
  
  - Be on the look out for emails from us this summer! (check your email!)
  
- For those on Exondys 51, you will be notified by your clinic if they are participating in the EVOLVE registry.
Conference Polling
Conference Polling Goals

- **Gain a better understanding** from families on a range of topics
- **Provide additional insights for discussions** taking place during conference
- **Inform drug development, trials, and care** through the lens of patients, caregivers, and families
Polling Guidelines

Polling will take place on Thursday, Friday, and Saturday in between sessions.

Those watching the stream can also participate.

To participate visit: pollev.com/ppmd

Wireless code: PPMD2019

Polling Questions

• Questions are for people with Duchenne and Becker, their caregivers and family members only.
• Some questions require only one response per family; we will indicate when we only need one response.
• Answers are anonymous.
Polling Advice

• Don’t fall victim to *peer pressure* with the answers on screen

  – Answer the polling question on your phone **before** looking at the results on the larger screen

• Don’t look at your neighbors responses
Polling warm up