Manipulating the Dark Side of Muscle Adaptation for Therapeutic Gain

Alan J Russell, PhD
Edgewise Therapeutics
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Forward-Looking Statements

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We have based these forward-looking statements largely on our current expectations and projections about future events and trends that we believe may affect our financial condition, results of operations, business strategy and financial needs. These forward-looking statements are subject to a number of risks, uncertainties and assumptions, including, among other things: negative impacts of the COVID-19 pandemic on Edgewise’s operations, including clinical trials; risks associated with the process of discovering, developing and commercializing drugs that are safe and effective for use as human therapeutics and operating as an early clinical stage company; Edgewise’s ability to develop, initiate or complete preclinical studies and clinical trials for, obtain approvals for and commercialize any of its product candidates; changes in Edgewise’s plans to develop and commercialize EDG-5506 or any other product candidates; the potential for clinical trials of EDG-5506 or any other product candidates to differ from preclinical, interim, preliminary, topline or expected results; Edgewise’s ability to enroll patients in its ongoing and future clinical trials; operating results and business generally; Edgewise’s ability to raise funding it will need to continue to pursue its business and product development plans; regulatory developments in the United States and foreign countries; Edgewise’s reliance on third parties, contract manufacturers and contract research organizations; Edgewise’s ability to obtain and maintain intellectual property protection for its product candidates; risks associated with access to capital and credit markets; the loss of key scientific or management personnel; competition in the industry in which Edgewise operates; Edgewise’s ability to develop a proprietary drug discovery platform to build a pipeline of product candidates; general economic and market conditions; and other risks. Information regarding the foregoing and additional risks may be found in the section entitled “Risk Factors” in documents that Edgewise files from time to time with the Securities and Exchange Commission. These risks are not exhaustive. New risk factors emerge from time to time, and it is not possible for our management to predict all risk factors, nor can we assess the impact of all factors on our business or the extent to which any factor, or combination of factors, may cause actual results to differ materially from those contained in, or implied by, any forward-looking statements. You should not rely upon forward-looking statements as predictions of future events. Although we believe that the expectations reflected in the forward-looking statements are reasonable, we cannot guarantee future results, levels of activity, performance or achievements. Except as required by law, we undertake no obligation to update publicly any forward-looking statements for any reason after the date of this presentation.

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When We Exercise, Our Muscles Get Bigger and Stronger

NYT – ‘How to get strong’

- Effect of 16 weeks resistance exercise (2/week) in healthy older men (65 yrs)
- Most metrics increased 30-50%
- Individual muscle fibers approx. 30% bigger

More Protein and More Muscle – Dual Pathways to Speed Adaptation

• Combination of protein synthesis/degradation and controlled muscle injury

**New protein**

12 Weeks Exercise Training in Younger and Older People

- **High Intensity Training**
- **Resistance Training**

- Membrane stress to activate stem cells

**Skeletal Muscle Adaptation to Exercise Training**

DNA → mRNA → Protein → Mitochondria → Hypertrophy

- Increased DNA
- Increased mRNA
- Increased Protein
- Increased Mitochondria
- Increased Hypertrophy

**Oxidation capacity**

**Muscle mass and strength**

Cell Metabolism 2017: 25, 581–592

Nat Rev Immunol 2017:17 p165
Adaptation Balance and Taking it Too Far!

Excessive or unaccustomed exercise

- Pain
- Stiffness
- Decreases in range of motion
- Increased muscle injury biomarkers (CK and other proteins)
Skeletal Muscle is Comprised of Slow (type I) and Fast (type II) fibers

Humans are ~50/50% fast/slow
Slow fibers are Less Prone to Disruption

- 30 mins controlled eccentric exercise
- Muscle biopsy taken immediately after exercise

MORE DISRUPTION IN FAST

MORE STEM CELL MOBILIZATION IN FAST


J Gerontology Ser A 64A:No. 3, 2009 p332–339
Do we have different fiber injury susceptibility to maximize adaptation but minimize the risk of disabling injury?

**SLOW FIBERS**

- Slow adaptation but less chance of injury?

**FAST FIBERS**

- Fast adaptation but greater chance of injury?
Becker Muscular Dystrophy (BMD) – Partially Functional Dystrophin

- 4,000-5,000 patients in the U.S.
- Later onset versus DMD, typically 8-15 yrs.
- Variable progression for mobility (late 30s) and cardiomyopathy
Normal Contraction Leads to Excessive Degeneration in Dystrophic Muscle

- Degeneration requires inflammation for repair
- Chronic inflammation leads to fibrosis
- Fibrosis leads to muscle loss and disability

Sue Brooks, Dennis Claflin, Sunny Yu, University of Michigan
Therapeutic interventions have focused on all elements of the degeneration process.

Approved therapies
- **High dose steroids** – alter inflammation to extend ambulation approx. 2 years but comes at a cost.
- **Antisense oligos** – weekly injections increase expression of shortened dystrophin in some patients. Unknown efficacy.

Developing therapies
- **Gene therapy** – delivery of a micro version of dystrophin via AAV virus. Inherent challenges with safety, efficiency of delivery and micro-dystrophin functionality.
Understanding the Function of Dystrophin – A Molecular Connector

Dystrophin connects contractile proteins to the membrane and surrounding matrix of fibers

With dystrophin – fibers support each other

No dystrophin – fibers contract without support
Similar to Healthy Muscle, Fast Fibers are More Sensitive in DMD

Fast Muscle Fibers Are Preferentially Affected in Duchenne Muscular Dystrophy

Cecelia Webster,*† Laura Silberstein,*†
Arthur P. Hayes,§ and Helen M. Blau*†
*Department of Pharmacology
Stanford University School of Medicine
Stanford, California 94305
§Department of Pathology
Division of Neuropathology
College of Physicians and Surgeons
of Columbia University
New York, New York 10032

Enrichment of muscle regeneration biomarkers in fast but not slow muscle fibers of young DMD kids

Black shading indicates eMHC-positive fibers

Regeneration Marker
Expressed mostly in fast but not slow fibers
Injury Biomarkers Tell the Same Story - Fast but not Slow Fiber Biomarkers are Elevated in BMD and DMD

A Left-Field Strategy to Rebalance Dystrophic Muscle?

Would selective inhibition of fast fiber contraction stop muscle degeneration but allow mixed fast/slow muscles to still function?
The Target – Fast Skeletal Muscle Myosin, the Engine of Force Generation

- A specialized motor protein that generates force by consuming ATP
- Different types of myosin motor are used by different types of ‘striated’ muscle (fast vs slow vs cardiac)
- *The motor in slow fibers and the heart is the same*
Identifying a Selective Fast Myosin Inhibitor

700,000 starting compounds

ATP consumption rates of homogenized native skeletal and cardiac muscle

Lead selective inhibitors

Synthesis and optimization of >1000 chemical derivatives of lead hits

EDG-5506!
EDG-5506 – A Potent, Selective Fast Skeletal Muscle Myosin Inhibitor

Selective inhibition of fast skeletal muscle suspension ATPase (but not slow/cardiac)

Concentration-dependent inhibition of force in isolated mouse muscle
EDG-5506 Stops Fiber Breakdown in Contracting DMD Muscle

Claflin, Su and Brooks. U Michigan

DMD muscle (mdx mouse) no treatment

DMD muscle (mdx mouse) 0.3 µM EDG-5506

Contracting at 100%

Contracting at 85%

mdx mouse lumbrical muscle – 20, 1 second maximal isometric contractions (video sped up)
EDG-5506 Reduces DMD Mouse CK After Exercise Testing without Altering Performance

Rotarod performance

Post-exercise plasma CK
(Injury marker)
EDG-5506 Decreases CK and Increases Activity in DMD Dogs

*Decreased injury biomarker (plasma CK)*

*Increased activity measured with an activity monitor*

![Graph showing decreased CK activity after dosing with EDG-5506](image1)

![Graph showing increased daily average activity after dosing with EDG-5506](image2)
Progressing EDG-5506 into Clinical trials

- Founded in Boulder, CO July 2017
- Oct 2017 – discovery work to identify promising compounds starts
- May 2018 – Discovery and advancement of EDG-5506
- Oct 2020 – Started dosing healthy volunteers
- Oct 2021 – Starting dosing adults with Becker muscular dystrophy
- Dec 2021 – Start of 1 year extension study in Becker
- 2H 2022 – Start of Becker phase 2 study
- 2H 2022 – Start of Duchenne muscular dystrophy studies
EDG-5506 Phase 1 Study Conducted in Healthy Volunteers and Participants with Becker Muscular Dystrophy

Trial Design

Phase 1a SAD in HVs*

Phase 1a MAD in HVs* (14 Days Daily Dosing)

Phase 1b BMD Cohort (14 Days Daily Dosing)

* All HV cohorts were randomized 3:1 active:placebo

Key Endpoints

Primary Endpoints

- Safety and tolerability at 20 mg over a 14-day period in BMD

Participants were monitored as inpatients for 16 days, with follow-up 1 and 4 weeks after completion of dosing.
### Participants in the BMD Phase 1b Had Significant Functional Impairment

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>BMD Participants (N=7)</th>
<th>Age Normative Values</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Age</strong></td>
<td>33.8 years</td>
<td></td>
</tr>
<tr>
<td><strong>Functional Measures (median)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10-meter walk/run</td>
<td>8.3 sec</td>
<td>&lt; 4 sec</td>
</tr>
<tr>
<td>Rise from floor</td>
<td>20 sec</td>
<td>&lt; 3 sec</td>
</tr>
<tr>
<td><strong>Serum Creatinine (mean, mg/dL)</strong></td>
<td>0.58</td>
<td>0.92 - 1.16</td>
</tr>
<tr>
<td><strong>Serum Creatine Kinase (mean CK, U/L)</strong></td>
<td>1,347</td>
<td>&lt;205</td>
</tr>
</tbody>
</table>

- Functional tests show significantly compromised or lost function
- Low creatinine consistent with decreased muscle mass
- Elevated CK levels reflect ongoing muscle damage
EDG-5506 was Well-Tolerated in BMD Subjects

<table>
<thead>
<tr>
<th>TEAE</th>
<th>Placebo N=2</th>
<th>EDG-5506 (20 mg) N=5</th>
<th>Total N=7</th>
</tr>
</thead>
<tbody>
<tr>
<td>Any TEAE</td>
<td>n (%)</td>
<td>n (%)</td>
<td>n (%)</td>
</tr>
<tr>
<td>Dizziness</td>
<td>2 (100%)</td>
<td>5 (100%)</td>
<td>7 (100%)</td>
</tr>
<tr>
<td>Euphoric mood</td>
<td>0</td>
<td>2 (40%)</td>
<td>2 (29%)</td>
</tr>
<tr>
<td>Musculoskeletal stiffness</td>
<td>0</td>
<td>2 (40%)</td>
<td>2 (29%)</td>
</tr>
<tr>
<td>Somnolence</td>
<td>0</td>
<td>2 (40%)</td>
<td>1 (14%)</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>0</td>
<td>1 (20%)</td>
<td>1 (14%)</td>
</tr>
<tr>
<td>Nausea</td>
<td>0</td>
<td>1 (20%)</td>
<td>1 (14%)</td>
</tr>
<tr>
<td>Fatigue</td>
<td>0</td>
<td>1 (20%)</td>
<td>1 (14%)</td>
</tr>
<tr>
<td>Vessel puncture site bruise</td>
<td>0</td>
<td>1 (20%)</td>
<td>1 (14%)</td>
</tr>
<tr>
<td>Back pain</td>
<td>0</td>
<td>1 (20%)</td>
<td>1 (14%)</td>
</tr>
<tr>
<td>Pain in jaw</td>
<td>0</td>
<td>1 (20%)</td>
<td>1 (14%)</td>
</tr>
<tr>
<td>Headache</td>
<td>0</td>
<td>1 (20%)</td>
<td>1 (14%)</td>
</tr>
<tr>
<td>Presyncope</td>
<td>0</td>
<td>1 (20%)</td>
<td>1 (14%)</td>
</tr>
<tr>
<td>Nasal congestion</td>
<td>0</td>
<td>1 (20%)</td>
<td>1 (14%)</td>
</tr>
<tr>
<td>AEs of special interest</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

- No change in grip strength
- All AEs were mild (Grade 1); AEs were transient and generally declined with increasing exposure
How to Detect an Early Activity Signal in BMD?

- Two weeks dosing enough time to assess short-term safety and tolerability but not long enough to measure beneficial effects on strength etc
- Before treatment with EDG-5506, BMD participants had elevated levels of CK as a result of ongoing muscle injury
- Were there other circulating proteins that also leak from muscle?
- Did short-term EDG-5506 stabilize muscle to reduce these proteins?
SOMAscan Aptamer Technology Enables Relative Measurement of 7000 Proteins for a Single Blood Sample

- ‘Slow Off-rate Modified Apatamer Scan’
- Aptamers are modified with a biotinylated photocleavable linker developed by Somalogic in Boulder, CO
- Aptamers are mixed with plasma samples then added to biotinylated beads
- Bead-bound proteins are biotinylated and beads are then washed and linker cleaved
- Protein/Aptamer complexes are bound to streptavidin beads and aptamers are then dissociated and quantified on a chip
- Note – SOMAscan optimized against human proteins, aptamers may not cross-react with proteins from other species
Using SOMAscan 7,000 Analyte Set, A Proteomic Signature for BMD was Identified

Baseline plasma samples (n=7) were compared to samples from healthy volunteers (n=25).

Proteins filtered by magnitude of difference (≥1.5X) and adjusted p value (<0.05) vs. HV.

Baseline analysis identified a fingerprint of 125 elevated proteins in BMD.
The Majority of Signature Proteins are Lowered by EDG-5506

Changes in BMD biomarkers vs. placebo (Day 1 vs Day 14)

BMD biomarkers responsive to increased exposure to EDG-5506

* EDG-5506 20 mg
Use of a Controlled Exercise Study to Annotate Our Becker Response Signature

- EDG-5506 lowered most proteins that were elevated in Becker vs healthy individuals
- However – what evidence do we have that any of them are related to exercise-induced injury?

Taking advantage of an ongoing collaboration with John Vissing (University of Copenhagen)
Defining a signature of proteins that directly increase with exercise in Becker

- Using controlled exercise, we established a set of 24 proteins that are elevated pre-exercise in Becker vs healthy and are then further elevated with exercise.
EDG-5506 Significantly Reduces proteins associated with injurious exercise in Becker muscular dystrophy

- Reduction of all annotated injury biomarkers with EDG-5505 in Becker muscular dystrophy
EDG-5506 Also Reduces proteins that are Independent of Exercise – Changing biomarkers outside of basic muscle protection?
Summary and thanks

• Building on learnings from muscle adaptation to exercise and how different skeletal muscle fiber populations respond, we’ve devised a novel approach to reduce muscle stress in two grave muscle diseases
• Early tolerability and biomarker data suggest that EDG-5506 has the potential to significant impact skeletal muscle health in Becker muscular dystrophy
• Studies start in Duchenne muscular dystrophy in 2022!

Co-Founders (Orbimed) – Peter Thompson, Badreddin Edris
Edgewise (Boulder, CO) - Mike DuVall, Ben Barthel, Ying Qian, Angela K. Peter, Breanne L. Newell-Stamper, Kevin Hunt, Stephen Schlachter, Ben Robertson, Behrad Derakhshan, Kevin Koch
University of Colorado, Boulder - Carlos Vera, Leslie A Leinwand
University of Michigan - Yu Su, Dennis R Claflin, Susan V Brooks
Texas A&M - Peter Nghiem, Alexis Rutledge
SOMAscan – Larry Gold, Luong Luu, Caylee Martens, Cole Zimmerman