

### February 2022

# Smith-Kingsmore Syndrome Foundation

### Rare Disease Day is Monday February 28th!

You can get involved in multiple ways to support Rare Disease Day:

- 1. Join the <u>SKS Rare WOD</u>. See the work-out below and visit this link <u>here</u> to learn more about how you can work-out to support SKS research
- 2. Go to <u>Twibbonize</u> to upload your Social media profile pic with SKS Rare Disease Day frame
- 3. Share your SKS Story with the SKS Foundation for our family story page. See the <a href="story page">story page</a> for examples and email us <a href="here">here</a>
- 4. Share your story with Nord. See this link here
- 5. Share your SKS Story on social media and use hashtags #RareDiseaseDay & #smithkingsmoresyndrome

All of these speak to our community mission of raising awareness of this ultra-rare disease!

\*click hyperlinks for more details



### **Current Research:**

- Dr. Liu, Dr. Prada & the Cincinnati Children's SKS Team have submitted their first publication supported by SKSF funded research. This will be a comprehensive paper on SKS & will open up new avenues for scientific inquiry
- Dr. Yang, the SKS post-doc at UF, research continues to evaluate the differences between 20 + different MTOR variants

### **Future Research:**

 Dr. Prada & Dr. Krueger, at Cincinnati Children's Hospital, are evaluating surveys collected and looking at brain function and behaviors. Look for the email from Lindsey E. Aschbacher-Smith requesting MRI and/or EEG data.

### Want to Enroll in SKS Studies?

• Email Lindsey at <u>Lindsey.Aschbacher-Smith@cchmc.org</u> for more details.

### Calling all German, Belgium & Netherland Families:

• Email Dr. Reijnders at margot.reijnders@mumc.nl for more information on the mTORopathy Expertise Center

### February 26th **Workout for Rare Disease Day**



Mission: See Families, Friends, and Communities are virtually coming together for the Well on February 26th. During a time of social distancing and quarantine this is your chance to celebrate by joining a cause worth working out for

Execution: CRF Athletes compete at various levels to complete the Sto 1997 Word. Sto Athletes come together (in spirit & virtually) in order to raise awareness for Rare Diseases Month and celebrate the first ever Sto West Word. Why such a hard workout? To "Be abnormal, among the normal." Just like these rare and extraordinary children.

#### 2008 BIRD WOD Athletic Levels:

Varsity: Athletes compete in all four prescribed workouts. There are no substitution lifts/movements at the varsity level. Do the workout as prescribed.

Junior Varsity: SE Athletes compete in two of the four West West events with athlete choice of rest time in-between.

Relay: We Athletes compete as a team to complete all events of the WALLAND Web.

Athlete Choice: Athletes pick any event to complete to the best of their abilities.



### **RARE WOD 2022**

It's time again for all the SKS athletes to sweat for SKS. Join us on for the world famous Rare WOD (workout of the day), powered by SKSF, on Saturday, February 26!

Click here to learn more about this event!

**Donate** today & support our SKS athletes!

### SKS RARE WOD GEAR FT: 5 Rounds

- -21x Double-Unders
- -15x Deadlift(Bodyweight)
- -9x Push-Press(1/2 Bodyweight)

#### MRAP: 15 min

- -21x Pull-Ups
- -15x Thrusters (1/2Bodyweight)
- -9x Toes to Bar

\*Each round completed = 1min off overall time.

FUN: 5k

### MOM (Ascending 1,2,3...)

- -Push-ups
- -Air-Squats
- -Burpees

\*Each minute completed = 1min off overall time.

# The Smith-Kingsmore Syndrome RECE Workout Of the Day



### RARE WALK

Dedicate a walk at any distance and pace to a child with a rare disease.

### SKS RARE WOD NO GEAR

### FT: 5 Rounds

- -21x Jumping Jacks
- -15x (eachleg) Lunges
- -9x Push Ups

### MRAP: 15min

- -21x Back Extensions
- -15x Burpees
- -9x V Ups

\*Each round completed = 1min off overall time.

Fun: 5k

### MOM (Ascending 1,2,3...)

- -Push-Up
- -Air Squat
- -Burpee
- \*Each minute completed = 1min off overall time.

### 2022 Foundation Initiatives

We continue to work with our dedicated SKS experts to raise awareness and increase research potential. Be on the lookout for these incredible initiatives:

- SKS Newly Diagnosed Informational Packet
- Up-To-Date SKS Report
- GeneReview SKS Report
- SKS NORD Report
- SKS Centers of Excellence where medical professionals are up to date on providing evidence based care to people living with SKS



# Smith-Kingsmore Syndrome Global Patient Registry Update February 2022

200+

Diagnosed with SKS Worldwide

69

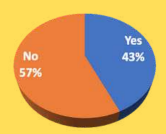
Participants from Australia, Belgium, Brazil, Canada, France, Italy, Japan, Kuwait, Mexico, Netherlands, New Zealand, Spain, UK, USA



64% Male 36% Female

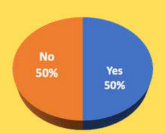
\* 44 participates have completed SKS specific questionnaire

# **Seizures**43% have had at least one seizure

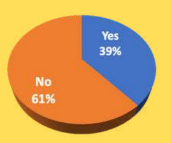


# Interception & Body Temp Dysregulation

50% Have difficulty feeling full/hungry, hot/cold, and/or overheats easily



# Hyperphagia 39%



# Neurodevelopmental & Behavior

