

“UBUNTU”

“More united and motivated to collaborate together”



(ITHACA's ePags - PAB carousel)





- Tomasz Grybek
tomek@fundacjabb.pl
- FBB - Foundation of Borys the Hero.
- True Partners in the rare disease world.



- Juan Antonio
admin@kbg syndrome.org
- Asociacion Espagnola Syndrome KBG
- Health is a state of complete physical, mental and social well-being and not merely the absence of disease or infirmity.



- Eszter Szabóné Katona
szabo.esther@gmail.com
- Disorder of the Corpus Callosum
Hungary Foundation
- Community with understanding



- Ellen Koekoeckx
ellen.koekoeckx@cureangelman.org
- FAST - Foundation for Angelman Syndrome Therapeutics
- Working together towards a better life for our loved ones living with a rare disease



- Daniele Palumbo
dp@champ1foundation.eu
- CHAMP1 Foundation – Europe
- If you don't try, you'll never succeed."



- Katarzyna Świeczkowska
katarzyna.swieczkowska@psoni.gda.pl
- Polish Association for Persons with Intellectual Disability.
- Developing networks, networks for development.



- Silvia de la Flor
delaforsilvia@gmail.com
- KAT6 France
- Syndrome KAT6A & Resilience



- Gerritjan Koekkoek
gerritjan@cdlsworld.org
- CdLS-World
- Cornelia de Lange syndrome
- Everyone knows something,
together we know everything



- Carole Herman
caroleheman2012@gmail.com
- ADNP France
- Hope with amis de ADNP



- Ana González Hernández
ana@ctnnb1-foundation.org
- Extra rare diseases
- Extraordinarily united



- Andrej Drdul
andrej.drdul@gmail.com
- Slovak Association for Spina Bifida and/ or Hydrocephalus
- IF - International Federation for Spina Bifida and/ or Hydrocephalus
- We transform fear from the unknown for friendship and trust. Our belief is that every person with a disability and their families as well may live a full-fledged, joyful life.



- Dorica Dan
dorica.dan@eurordis.org
- Prader Willi Association, Romanian
National Alliance for Rare Diseases
- Living together, learning together!



- Benoit Fourcroy
benoit.f@spina-bifida.org
- ASBH - spina-bifida.org
- Advances in healthcare and peer support



- Erika Stariha
erika.stariha@satb2europe.org
- SATB2 Europe
- Hope Becomes real when we come together



- Nina Knight
ninasknight@hotmail.co.uk
- Acrodysostosis Support & Research UK World
- Together we accelerate the science and create a pathway to treatments



- Sue Routledge
sue@pitthopkins.org.uk
- Pitt Hopkins UK and Stichting
- Pitt-Hopkins syndrome
- Learning and going forward together



- Ioel Detton
assonoonan@gmail.com
- Noonan France
- Alone, we go faster; together, we go further



- Vesna Vujičić
22qexyu@gmail.com
- 22q ex Yu, Serbia
- Together we can change the future



- Anne Lawlor
a.enekwe@gmail.com
- 22q11 Ireland
- Together for Rare Families



- Antje Enekwe
a.enekwe@gmail.com
- SLO Deutschland e.V, Smith-Lemli-Opitz Syndrome
- Let us make a difference for people with rare diseases.



- Olga Sophocleous
anefeli@cytanet.com.cy
- Angelman syndrome Greece
- Common vision for our rare patient's better future



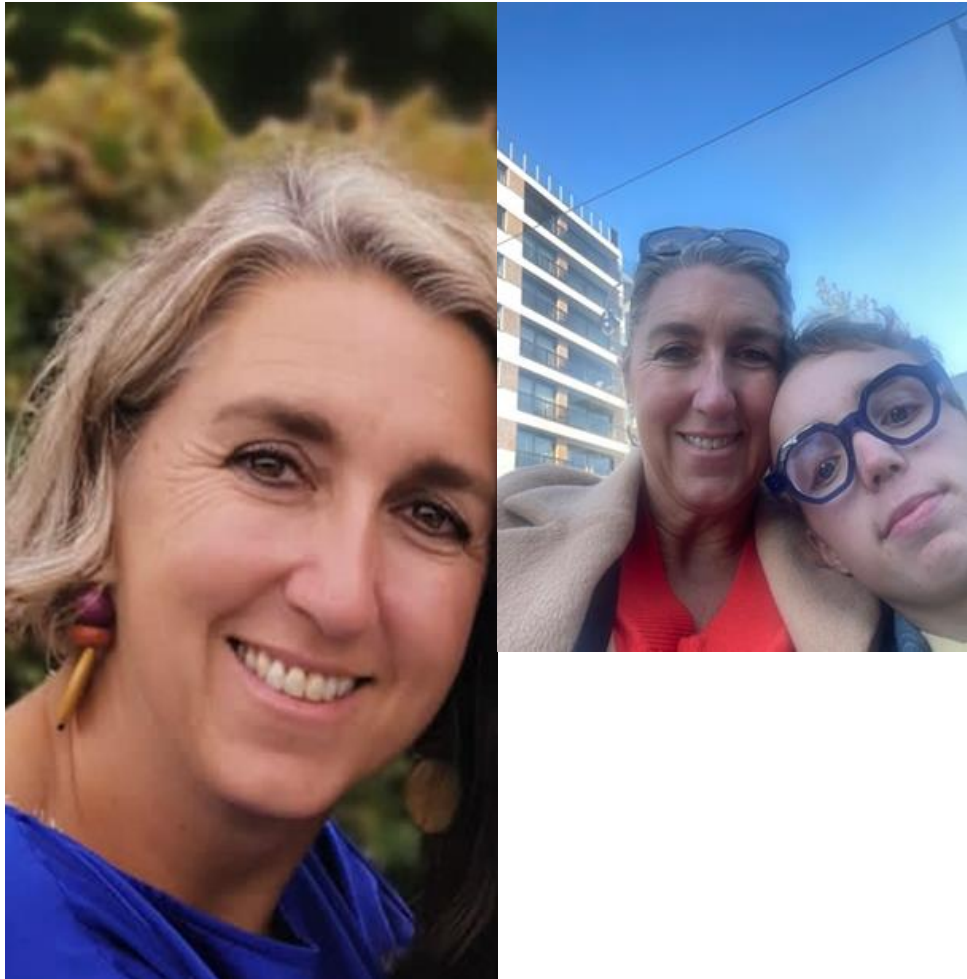
- Tanja Zahlten
tahischle@hotmail.com
- German Bundesverband WBS
- Williams Beuren Syndrom
- Share knowledge and learn from each other



- Ines Fernandez Ulibarri
ifernaul80@hotmail.com
- Asociación Kleesfstra España
- Kleefstra syndrome
- Together to Change the Future of Rare



- Annalisa Scopinaro
annalisa.scopinaro@yahoo.it
- APW Italia
- Williams Syndrome
- Together we can



- Kelly Verbruggen
adnpkids.hvdasbelgium@gmail.com
- ADNP kids Belgium
- Helsmoortel - Van der Aa syndrome
(aka ADNPsyndrome)
- Striving to a better understanding of
Helsmoortel Van der Aa syndrome
and supporting parents across the
world



- Rui Barbosa Guedes
rui@phelan-mcdermid.pt
 - APMP Associação, Portugal
 - Phelan-McDermid
-
- Beacon of Hope for brighter future



- Monica Bertoletti
monica@aibws.org
- AIBWS Odv Italia, Association
- Beckwith-Wiedemann Syndrome
- Sharing, collaboration, better perspective



- Tanja Zdolšek Draksler
tanja.zdolsek@ijs.si
- IDefine Europe Foundation
- Kleefstra syndrome
- Data is the key



- Pogány Gábor
pogany@williams.org.hu
- Hungarian Williams Syndrome Association (HWSA), (FEWS) – (HUFERDIS, Rare Diseases Hungary)
- Williams Syndrome
- Start by doing what's necessary then do what's possible and suddenly you are doing the impossible” (Saint Francis of Assisi)



- Jessica McAndrew
jessica@kabukisynndromefoundation.org
- Kabuki Syndrome Foundation
- Building brighter futures for those affected by rare disease



- Marta Balula
marta.balula@raramente.org
- Raramente, portugal
- Rare syndromes
- Empowering to advance Care, Knowledge, and Advocacy for Rare Lives



- Ammi Andersson
ammi.andersson@cordnode.se
- RBU Rörelsehindrade Barn och Ungdomar Stockholm
- Spina Bifida and Spinal Dyphrasism
- Cooperate with clinicians to create an equal specialist care for rare conditions in every country within EU

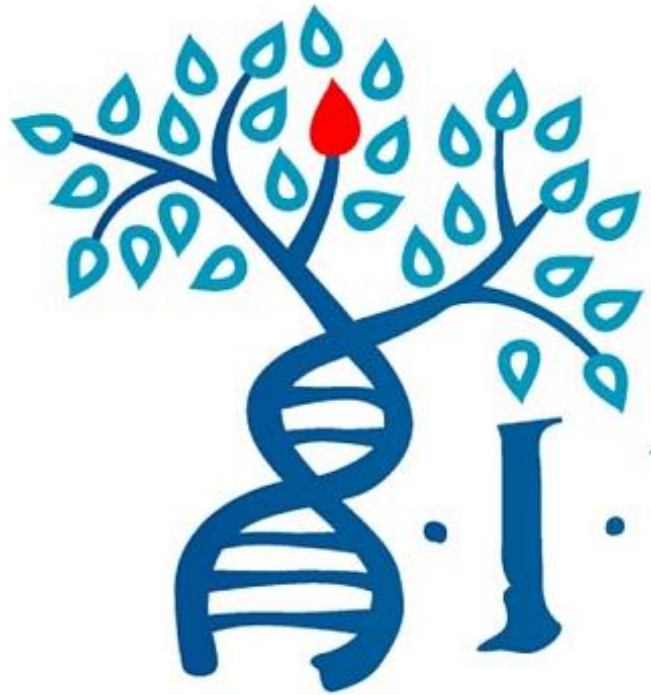


- Lyndsey Walsh
lyndsey.1@hotmail.com
- Rare Diseases Ireland, 22q11
Ireland and Ultra-Rare Diseases
- Together we are a strong
partnership for change



- Suzàn Genova
suzan.m.genova@gmail.com
- FAST Bulgaria - Angelman syndrome; Nataliya Foundation - Isovaleric acidemia
- A shared path to hope

Thank you for your attention !



UBUNTU
"I am because we are."

SI·THA·KA