Delivering on Canada's Rare Disease Investment



Rare Disease Day Conference

March 28 – 29, 2023

Ottawa Marriott

100 Kent Street, Ottawa

*Virtual option available



Tuesday, March 28, 2023 Rare Disease Infrastructure Essential to Optimal Drug Benefit	
8:30 AM - 9:00 AM	Registration & Continental Breakfast Victoria Ballroom (2 nd Floor)
9:00 AM - 9:25 AM	Welcome and Opening Address Durhane Wong-Rieger, CORD Keynote Speaker: Canada's Commitment to Rare Disease Patients and Families • Pamela Aung-Tin, Assistant Deputy Minister, Health Products and Food Branch, Health Canada Keynote Panel: A Vision for Rare Disease in Canada • Alex Munter, Children's Hospital of Eastern Ontario (5 min) • Brigitte Nolet, Roche (5 min)
9:25 AM - 9:35 AM	Challenges to Address Rare Patient Experience in Canada Feedback from patients and families to CORD survey on navigating the healthcare system: what is working, not working, and needs to be changed. What should we do with \$1 billion to improve lives of patients and families with rare disease? • Hyejin Park, Ipsos (10 min)
9:35 AM - 9:45 AM	Development of Canada's Rare Disease Network: Centres of Expertise Setting the Stage • Daniel MacDonald, Health Canada (5 min) • Ian Stedman, York University (5 min)
9:45 AM - 10:00 AM	Interactive Discussion with Expert Panel and Participants Three brief presentations will set the stage for implementing Canada's Rare Disease Network of Centres of Expertise as fundamental to optimal patient care and essential to optimal drug access and value. Presenters: Durhane Wong-Rieger, CORD Leanne Ward, CHEO François Bernier, University of Calgary

- **Centres of Expertise:** How can the Canadian Rare Disease Network assure that every patient and family affected by rare disease gets timely diagnosis, access to specialists, and integrated comprehensive care? How does the CRDN function within a WHO Global Rare Disease Network? Durhane Wong-Rieger, CORD (5 min)
- **Specialized Network:** What is an example of a specialized network that serves a specific disease cluster, how it is organized, who are involved, and what services and supports are offered? What additional resources are needed to expand and sustain these specialized networks within a Canadian Rare Disease Network? Leanne Ward, CHEO (5 min)
- **Clinical Reference Centres of Expertise:** How can existing children's healthcare centres be organized to serve as the "backbone" of national Rare Disease Network and also to serve as the "hub" for support to local health providers? François Bernier, University of Calgary (5 min)

10:00 AM - 10:45 AM | Facilitated discussion: How to Optimize Rare Disease Drug Strategy

Leads:

- Bill Dempster, 3Sixty Public Affairs
- Durhane Wong-Rieger, CORD

Expert Panelists:

Sunita Venkateswaran, London Health Sciences Centre; Pranesh Chakraboty, CHEO; Riyad Elbard, Thalassemia Foundation of Canada;

François Bolduc, University of Alberta; Kym Boycott, CHEO; Kasha Mitton, Defeat Duchenne Canada; Cathy Evanochko, Tuberous Sclerosis Canada;

lan Stedman, York University; Kim Steele, Cystic Fibrosis Canada; Leanne Ward, CHEO;

Open discussion with panelists and all audience

1. The proposed national plan identifies key components, objectives and four "strategic" pillars; are these sufficiently comprehensive; if not, what is missing? 2. How does announced Rare Disease Drug Plan align with CORD's Rare Disease Strategy, which is based on networked Centres of Expertise, Value-based Managed Access to

Therapies, and Investment in Research?

- 3. What are the opportunities and risks inherent in the proposed national Drug Plan to advance diagnosis, care and treatment for patients and families? How can we leverage the opportunities and neutralize or minimize the risks?
- 4. What are the success factors to optimize benefits to patients? Who are other key players that need to be at the table to help design the plan and implement the strategy? What other investments and supportive mechanisms are needed, and how do we secure these?

10:45 AM - 11:00 AM Refreshment Break

11:00 AM - 12:00 PM | Case Studies with Expert Panel

The next are very brief overviews of case examples drawn from "real life" Canadian patient journeys that demonstrate the challenges, as well as facilitators, in the healthcare system.

- Muckle Wells/CAPS: "Diagnosis became a family affair." After over six decades of searching, this woman's entire family finally received its rare disease diagnosis when her granddaughter was born with some familiar symptoms. What helped was that a new treatment meant specialists were now alerted to the disease spectrum.
- 2. **Fabry Disease:** "Can multi-generation screening unlock the family tree?"

My father died at 29 years when I was four, and his brothers never made it to the age of 30; the women lived longer, to about 50 years. No one knew what killing them. Until at 52 years, I presented with heart problems and my cardiologist said, "Let's do genetic testing." I

- got a name for the disease, a new treatment, and a new chance for life."
- 3. **Aplastic anemia and PNH:** 'Lightning can strike twice."

 At 26 years old, I thought I would finally put my struggles behind me when I found a donor match for a life-changing bone marrow transplant. Until just before the procedure, they found I also had paroxysmal nocturnal hemoglobinuria (PNH), an ultra-rare condition that killed off healthy cells, including my new bone marrow cells.
- 4. **Myasthenia gravis:** "It's all in your head."

 I was in my late 20s when I started to have difficulty speaking. Every doctor said it was in my head. After my first child was born, things got worse but still no one believed me. What saved me was a sinus infection and my throat closed up, so I ended up in the ER, where a doctor gave me an injection and a diagnosis.
- 5. **Hereditary angioedema:** "Rare when remote is twice as hard." Because I live in rural Manitoba, I had to travel long distances to meet with allergists and immunologists. My symptoms with swelling that would subside for a while and then come back worse than ever didn't match with anything I had seen from years working in my local ER. Even with a diagnosis, the anxiety of an attack is always in the back of my mind.
- Cholangiocarcinoma: "Delayed diagnosis made a bad cancer worse."
 - I was happy and healthy. Then, within two weeks, I found out I had stage four bile duct cancer, and I was being scheduled for radiation and chemo. There are no good cancers but this is a particularly bad one.
- 7. **Spinal Muscular Atrophy:** Will a one-month delay make a lifetime difference?"
 - It was nearly a month before our son would be diagnosed with the most severe form of SMA, a rare genetic disease that weakens and wastes muscles. If he had been born in Ontario, he would have been tested at birth with a heel prick and had immediate access to treatment before symptoms set in.

Expert Panel Discussion

Expert panels will discuss case examples of patient journey challenges to diagnosis and access to specialists to identify what is working in the current system, what are the challenges experienced,

12:00 PM - 1:00 PM	 and what changes could be addressed in a Rare Disease Network model. What are the barriers and facilitators to timely diagnosis? What are the challenges and opportunities in access to specialists? How is access to comprehensive integrated multidisciplinary and supportive care for rare diseases working, or not? Lunch
1:00 PM - 1:15 PM	Value-based Healthcare and Real-World Evidence: TOWWERS Showcase Bridging Centres of Expertise and Managed Access to Therapies - Julie Frappier, Data 4 Actions
1:00 PM - 1:30 PM	Alternative Pathways for Access to Rare Disease Drugs: Brief Presentations and Facilitated Expert Panel Discussion • Managed Access/Value-Based Drug Access Schemes: Global Perspectives: What are different models of Managed Access Schemes and where and how are these being used in various countries to provide access to rare disease therapies? What are the benefits and risks of each approach? What are requirements for effective implementation, including risk-sharing financial arrangements, real-world evidence, and outcomes assessment. Fred Little, Pfizer Canada • Managed Access Schemes—Canadian Experience: What has been the Canadian experience with Managed Access Schemes? Where and how have they worked and what have been challenges to making them work? Tara Cowling, Medlior • HTA and Managed Access: How can health technology assessment contribute to the development of managed access plans? When and how should Canada's HTA agencies participate in MAS for rare disease drugs? Sylvie Bouchard, INESSS • Provincial Perspectives on Value-Based Rare Drug Access: How can a value-based approach to a rare drug access provide timely access to those who could benefit while also assuring affordability and sustainability in public drug plans? How can strategies for risk-sharing between pharma and drug plans accelerate access for patients and mitigate risk to the payers? Mitch Moneo, BC Ministry of Health

	Moderators:
	 Alexandra Chambers, Novartis
	Bill Dempster, 3Sixty Public Affairs
	Panelists:
	Fred Little, Pfizer Canada
	Allison Wills, 20Sense
	Tara Cowling, Medlior
	Sylvie Bouchard, INESSS
	Mitch Moneo, BC Ministry of Health
	Gaby Bourbara, Alexion Canada
	Rosalie Wyonch, CD Howe Institute
	Julie Frappier, Data 4 Actions
	Peter Brenders, BeiGene
2:30 PM - 2:45 PM	Afternoon Break
2:45 PM - 3:45 PM	Case Studies with Expert Panels
	Addressing Challenges to Access to Treatment through
	Managed Access Programs
	Cases: Access to Therapy
	1. SMA Type3: Access beyond clinical trials
	The treatment is here, it has been approved, but it's not available to
	me because of my age and my SMA type.
	2. Retinal Blindness: When there are no more cones, cannot treat
	Raising awareness through advocacy is the way we get our message
	heard and bring about real change.
	3. Fibrodysplasia Ossificans Progressive : Protect against future
	injury
	It's so tough as a toonager when you're finally ready to be more
	It's so tough as a teenager, when you're finally ready to be more
	independent and to start doing more on your own, to instead find
	independent and to start doing more on your own, to instead find
	independent and to start doing more on your own, to instead find yourself going backwards.
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	 independent and to start doing more on your own, to instead find yourself going backwards. 4. XLH: Clinical trials: an adventure far from home Once we had the diagnosis, that's when the real journey started — the around-the-clock phosphate supplements, the surgeries, the

- What is it worth for one more month to hug your kid? One more week? One more day? Most Canadians with ALS are being denied access because of very restrictive criteria.
- 6. **aTTP:** Rescue and prevent future crises

 You never know when a life-threatening episode will occur, so even in remission, the disease remains an ongoing challenge.
- 7. **LC-HAD/LC-FAOD:** Risk of bias ... statistical uncertainty I've been trying to avoid every little sickness for my whole life. The one thing I haven't been able to control, though, is the vision loss. They said no to the only effective treatment.
- 8. **VHL:** "The tumours are like salt and pepper all over the cord." With each spinal cord surgery, the scar tissue becomes more extensive.... I'm terrified they will grow and require more surgery. I can't keep doing this ...

Panel A: Brad Alyward, CORD; Sylvie Bouchard, INESSS; Claudia Caminit, Answering TTP; Wayne Critchley, Global Public Affairs; Chad Mitchell, Associate Deputy Minister, Alberta; Beth Vanstone, CF Get Loud (former)

Panel B: Alexandra Chambers, Novartis; Carrie Connell, Canadian FOP Network; Tara Cowling, Medlior; Jida El Hajjar, Loeys-Dietz Syndrome Foundation; Bob McLay, Sobi; Stephen Parrott, Canadian VHL Alliance

Panel C: Martine Elias, Myeloma Canada; Cheryl Greenberg, Children's Hospital Research Institute of Manitoba; Fred Horne, former Health Minister Alberta; Fred Little, Pfizer Canada; Mitch Moneo, BC Ministry of Health; Christine Mossa, Ipsen; Susi Vander Wyk, CureSMA Canada; Rosalie Wyonch, C.D. Howe Institute

Expert Panel Discussion

Expert panels will discuss case examples of patient experiences of challenges to timely access to appropriate treatments, including innovative medicines, to identify what is working in the current system, what are the challenges experienced, and what changes could be addressed with Alternative Pathways, including Managed Access Programs

	 What are best practices in other jurisdictions that should be considered for Canada? Clinical trials for rare/ultra-rare disease drugs are often small, short, and single-arm. Outcomes data may rely on surrogate measures, including biomarkers, and short-term patient-reported outcome measures (PROMs) that are not easily captured by conventional, validated, quantitative Quality of Life (QoL) scales. Because of the small patient population, cost may be high on a per-patient basis but have limited budget impact. How should regulators, value assessors, and payers take all of these factors into consideration when deciding on approval and access?
3:45 PM - 4:00 PM	Day 1 Wrap-up
5:00 PM - 7:00 PM	Networking Reception – Summit (29 th Floor)
	CORD is delighted to host an event for all conference participants to
	network with CORD Board of Directors, speakers, industry and
	fellow rare disease advocates.

Wednesday, March 29, 2023 Rare Disease Infrastructure Essential to Optimal Drug Benefit	
8:30 AM - 9:00 AM	Continental Breakfast Victoria Ballroom (2 nd Floor)
9:00 AM - 9:15 AM	Day 1 Recap Durhane Wong-Rieger, CORD
9:15 AM - 10:45 AM	Canadian Rare Disease Research in Drug Development, Trials, and Outcomes
	Interactive Discussion with Expert Panel and Participants Creating Canada's Rare Disease Research Network
	Canada boasts world-class researchers engaging in leading-edge innovative research programs focused on rare disease and precision drug development, including cell and gene therapy. We are also experiencing renewed and expanded investment by pharmaceutical companies in Canadian-based new drug development, clinical trials, and patient support programs for rare diseases. Finally, public and private investments are creating new initiatives and opportunities to partner with Canadian rare disease patients and patient organizations to ensure Canadians benefit from new knowledge and new technologies. Experts will discuss research initiatives and what these mean to rare disease patients and families and beyond. They will also discuss the benefits of creating a Canadian Rare Disease Research Network and the requisite requirements.
	Presenters, Panelists and Group Leads:
	Étienne Richer, CIHR, Institute of Genetics; Carrie McElroy, Sanofi Canada/RAREi; Thierry Lacaze, MICYRN; Risini Weeratna, National Research Canada; Stéphanie Michaud, BioCanRx; Sapna Mahajan, Genome Canada; Farah Bendahmane, Montreal InVivo; Susan Marlin, Clinical Trials Ontario, Brian Ballios, Toronto Western Hospital; François Bolduc, University of Alberta; Conor Douglas, York University

Presentations (10 min each)

- Social Pharmaceutical Innovation: Made-in and for Canada initiative for research and development of sustainable, accessible, and affordable pharmaceutical innovations to benefit patients and families, in Canada and beyond - Conor Douglas, York University
- Canadian Institutes of Health Research: What are current funded research projects that are focused on or have application to rare disease? What are future potential opportunities for research funding? What programs exist for training and supporting patients to engage in CIHR-funded initiatives? - Étienne Richer, CIHR, Institute of Genetics
- National Research Canada: What are initiatives led by, funded by, or supported by NRC that have benefit and relevance to rare disease patients in Canada, and beyond? What are specific aspects of the Canadian scientific, health and social ecosystems that make Canadian an effective and competitive environment for rare disease, precision, and targeted research and development? Risini Weeratna, National Research Canada
- Genome Canada: How has Genome Canada been active in the funding and support of research initiatives that benefit and/or support rare disease researchers, clinicians, and policy makers with direct and relevant benefits for patients and families? Sapna Mahajan, Genome Canada
- MICYRN: How does MICYRN function to attract, coordinate, stimulate, and support rare disease research and product development? What is the vision of MICYRN in the proposed Rare Disease Network and Rare Disease Drug Strategy? Thierry Lacaze, MICYRN
- Clinical Trials Ontario: What are the initiatives and capacities at CTO that directly and indirectly support researchers and clinicians in the recruitment and implementation of clinical trials and related activities, with direct engagement of and benefit to rare disease patients and families? Susan Marlin, Clinical Trials Ontario
- **InViVo Montreal:** How can this new tool provide a "scientific watch" on new therapies with potential indications in rare

	 conditions, including cell and gene therapies? The goal of this tool is to help payors, HTAs and the community at large to get a better sense of what's coming next and how to prepare. UHN Clinical/Research Project on Adult Retinal Disease: Unique pilot project to bridge gap for adult patients with retinal disease. Brian Ballios, Toronto Western Hospital Al and Rare Disease Research: Applications of Artificial Intelligence (AI) to understanding and advancing rare disease research with focus on Fragile X and autism. François Bolduc, University of Alberta RAREI (Canadian Forum for Rare Disease Innovators): informal network of research-based bio pharmaceutical innovators committed to monitoring, responding and shaping policy issues in the Canadian rare disease environment. Carrie McElroy, Sanofi
10:45 AM - 11:00 AM	Refreshment Break
11:00 AM - 12:00 PM	Expert Panel Discussion: Steps Toward Creating Canada's Rare
	Disease Research Network
	In what ways might a Canadian Rare Disease Research Network
	stimulate, support, and deliver on the following issues:
	R&D on Innovative Therapies
	Patient Outcomes Research
	Understanding Rare Diseases and Patient Histories
12:00 PM - 1:00 PM	Lunch
1:00 PM - 3:00 PM	Critical Success Factors? Improve research on rare diseases
1:00 PM - 1:40 PM	Patient Data Platforms - Brad Milson, IQVIA
	To support an effective Rare Disease Drug Strategy based on the
	three pillars of Centres of Expertise, Managed Therapeutic Access
	Schemes, and Research, an integrated data infrastructure is
	essential This would have the following objectives:
	(1) Integrate data from multiple stakeholders;
	(2) Expedite diagnosis and treatment;
	(3) Improve research on rare diseases;
	(4) Enable cost-sharing with multiple players;
	The infrastructure collects and links data flows from multiple
	sources and users sustainably via a standardized data structure and
	content, with the following core components:

(1	1) Multifunction patient portal;
(2	2) Electronic health records;
(3	3) Patient registries, especially for clinical trials;
(4	4) real-world data;
(5	5) Patient profiles of different patient groups;
(6	6) Standardize structure and language to facilitate a data sharing
a	cross domains and efficient data management and analysis.
1:40 PM - 2:20 PM E	conomics of Returning Value for Investment - Lindy Forte,
E-	eversana
R	Rare diseases are often severe, chronic, progressively debilitating
a	and life-threatening conditions with psychological and physical
e	effects that seriously compromise quality of life. It is important to
th	he understand the economic, social and quality of life impacts on
	ndividuals, families, and society, including medical and non-medical
	osts as well as loss of productivity, informal support, and
	systematic inefficiencies in diagnosis, care and treatment, with a
-	goal of establishing a potential return on investment for a properly
	designed and implemented integrated comprehensive rare disease
	system.
	Genomic Sequencing: Equitable and Accessible for Clinical
	Diagnosis - Magda Price, CHEO
	An update on the Genome Canada-funded AllforOne Health Data
	Ecosystem will be shared. This national data sharing initiative for
	are disease is proposed to support high quality clinical genomic
	esting as well as make some of this data available for research. The
	AllforOne Clinical Network will enable comprehensive variant
	haring between Canadian diagnostic laboratories for improvement
	of clinical genome-wide sequencing. AllforOne Connect will be a
	entralized registry to connect individuals with a rare disease to
	esearch studies they may be eligible for. This initiative will
	ultimately improve the health and wellness of Canadians with rare
	disease, be one an innovative contribution to rare disease research
	n Canada, and lay the foundation for precision health in Canada.
II	in Cariada, and lay the Touridation for precision health in Cariada.
3:00 PM - 3:30 PM	Next Steps

Conference Sponsors

The Canadian Organization for Rare Disorders acknowledges the contribution of all our Corporate Partners to improving the lives of patients and families with rare disorders.

We are especially grateful to the following that have supported this conference.























































