

# PxP 2025 SEPTEMBER 9-12

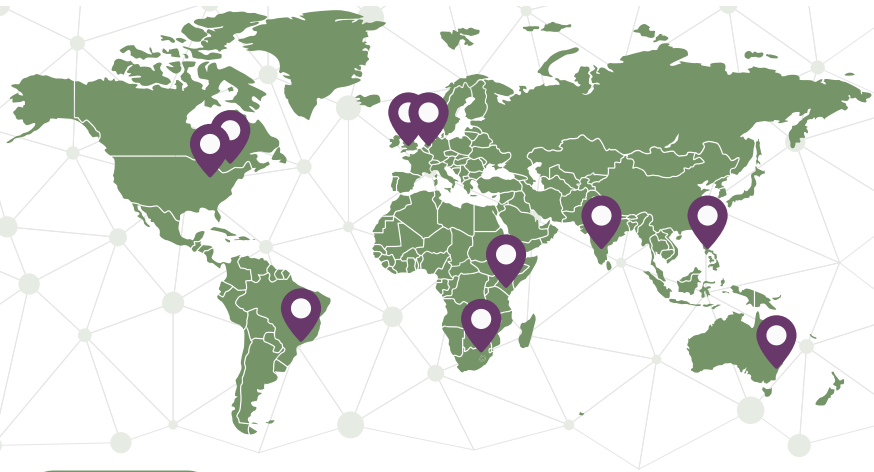
## VIRTUAL CONFERENCE

Join us in September for our free, virtual conference to share resources, mentorship and community with patient partners and others involved in health research. We hope that together, we can **plant the seeds** to promote patient engagement best practices and partner to make research stronger.



### What is PxP?

Standing for 'For Patients, By Patients', PxP is a patient-led initiative all about patients engaging as partners on health research teams. The annual, virtual PxP conference is a highlight of the initiative which launched in September 2023.

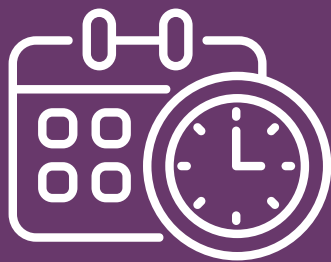


PxP 2025 is entirely led by an **international steering committee**, with members identifying as patient and public partners and is a Patients Included™ event.

# About the PxP Conference

**PxP is a must-attend event for these three reasons:**

1. PxP is specifically For Patients. While there has been excellent progress in the available patient engagement resources, many of these are designed for other members of the research team. Although we welcome health researchers to join us at the event in September, our key priority will be patient partners and other people with lived experience.
2. We are led entirely By Patients. Every topic and speaker on the PxP conference program has been chosen by our experienced international committee of patient partners.
3. PxP is about patient engagement in research. Many conferences focus on a specific research area, and have some patient engagement in them, but PxP focusses on how to do patient engagement and the experiences related to this.



For those based in Europe, Africa and America, the dates are September 9-11, and for those in parts of Asia and Oceania the dates are September 10-12. There will be core conference hours each day where content will be hosted live. New for this year, there will also be 'watch parties' for some of the sessions outside of the core hours. These watch parties mean that content will be played back for different time-zones and you can join those with others to chat in real-time.

Conference timings are available in a table at the bottom of this web page:

<https://pxphub.org/event/free-tickets/>



# Impact from past events

## PxP 2023

The first ever PxP conference took place on September 12–14, 2023. PxP 2023 was a three-day virtual conference that focused on boosting skills in patient engagement, trailblazing new approaches, and strengthening patient partnerships by raising the benchmarks for best practice.

**400+** registrants

from

**30+**

countries



**30+ speakers**

presented to attendees who self-identify as:

- Patient partner/Individual with lived experience
- Researcher
- Caregiver/Relative
- Clinician
- Trainee/Student
- Interested in learning more about patient engagement

## PxP 2024

PxP 2024 followed on September 10–13, 2024.

The event continued the conversation by valuing your voice as a patient partner, tackling the challenging topics about patient engagement in research and advancing best practice in research partnerships.

**1000+**  
registrants

from

**35+** countries



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*As a new and inexperienced patient partner, it was eye-opening to me. I learned to be much more careful about the emotional labor I offer in my interactions with studies and feel more justified in receiving financial exchange for doing so.” –PxP 2024 Attendee*

Learn more at [pxphub.org](https://pxphub.org).



# PXP Program 2025

## Day 1 What's happening and what's possible

Global voices, local impact: patient engagement in research around the world

Roles in research: stories of what these look like and offer

Journeys of impact: becoming and growing as a patient partner

Ask Me Anything: patient engagement in research in a changing world

## Day 2 Setting up for success and inspiring others

Laying the groundwork: setting up for success in patient partnership

Owning your lived experience: patient partner as part of your identity

Stronger together: building capacity on research teams

Ask Me Anything about research or about how I engage patients in my research

## Day 3 Pushing the envelope

At the cutting edge: innovations in patient engagement

Patients leading the way: research from within

More than a paper: ways to share research

You Can't Ask That: the future of patient engagement in research

