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Inspired by chance: valuing patients’ informal contributions to research

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Serendipitous contributions from patients that influence the research agenda should be better recognised and acknowledged, argue Sebastian Crutch and colleagues

Public involvement is increasingly considered a required part of health research. Typically defined as research being carried out “with” or “by” members of the public rather than “to,” “about,” or “for” them,1 public involvement is different from engagement (provision and dissemination of information and knowledge about research) and participation (taking part in a research study). Typical activities include identifying research priorities,2 choosing and co-designing outcome measures,3 advisory group membership, reviewing research materials, and carrying out the research.

Several major journals and funders have adopted strategies for enabling patients and the public to shape their aims and ways of working. Some also require researchers to make explicit statements about how—and importantly when—that involvement in the research process has occurred (eg, The BMJ’s patient and public partnership campaign4). Other recent steps to improve standard practice include guidelines for researchers and the public,5 6 the development of public involvement standards,7 and suggested reporting guidelines (GRIPP2) for public involvement in research in research.8 These changes sit within a broader landscape of calls for greater involvement of patients and the public in scientific conferences,9 and
autonomous patient-led movements to lead and crowdfund research\textsuperscript{10,11} and innovate systems of care and support.\textsuperscript{12}

A plethora of involvement frameworks have been developed.\textsuperscript{13} However, formal frameworks may not always capture the richness and variety of informal contributions to research that occur serendipitously, retrospectively, or independently. These informal contributions may originate in everyday conversations and interactions between researchers and people living with health conditions, including those with cognitive impairments that may make formal meetings and documents less appropriate.

**Inspiring new research and challenging assumptions**

When people with lived experience of dementia share their questions, experiences, and uncertainties, it may not be with any express intent to contribute to research. But these insights often have the effect of inspiring research by generating new ideas or hypotheses in the mind of the listener (or someone else with whom the gist of the conversation is later shared), challenging existing assumptions about their condition, or posing questions that can be investigated through academic research. Such conversations may take place during clinical interviews, at support group meetings, by letter or email, or over a cup of tea between research assessments. Critically, these inspiring contributions may occur at any time, not just conveniently during the planning phase of a project, and they might lead to new research activity immediately or some time later. One of the most striking examples of this we have known led to the discovery of the first gene for Alzheimer’s (box 1).

**Box 1: Family connections**

Carol Jennings’ father was diagnosed with Alzheimer’s disease in his mid-50s. Subsequently an aunt and uncle began to experience symptoms, and she recalled her grandfather having similar problems. Describing events in 2010, Carol said: “I put two and two together and thought there’s something funny going on here. This is the same family, developing the same symptoms, at a similar age … I made a family tree, I found out lots of names and I thought this was interesting to someone who could do something about it.” Carol sent this information to MR’s team and had an important role in the ensuing research, gathering together her whole family on the paternal side for blood tests. This research led to the discovery of the first gene for Alzheimer’s disease.\textsuperscript{14}

Open ended conversations with people with lived experience of a condition can also provide an opportunity to reflect on and question the status quo in a given area, and to challenge assumptions and standard practice. At a posterior cortical atrophy (PCA) support group meeting in London in March 2009, one member (co-author Simon Rosser) raised a simple but critical point that the use of different terminology for the condition (PCA, visual variant Alzheimer’s disease, biperiatal Alzheimer’s disease, Benson’s syndrome) resulted in a
lack of understanding among patients, carers, and professionals. His comment sparked the formation of an international working party and the first multicentre research diagnostic criteria for PCA.\textsuperscript{15}

The sharing of inspirational experiences is not limited to the spoken or written word, as we discovered when we had the privilege of working with William Utermohlen, a renowned artist primarily of portraits and murals. In 1995, he was diagnosed with Alzheimer’s disease, and over the following five years created a series of self-portraits that altered strikingly in their style and representation of perceptual form and spatial relationship (fig 1).\textsuperscript{16} His work continues to communicate something about both the experience of living with a dementia and the biology of Alzheimer’s disease that is more powerful than any brain scan, blood test, or scientific graph.

Posthumously, his comments and descriptions, and the lasting effect he had on clinicians and researchers, inspired the Created Out of Mind residency (2016-18) at The Hub, Wellcome Collection, which aimed to explore, challenge, and shape perceptions and understanding of dementias through science and the creative arts.\textsuperscript{17}

\textbf{Fig 1} William Utermohlen, Self Portrait with Saw 1997, 35.5 x 35.5 cm, Boïcos Collection, Paris. Courtesy of Chris Boïcos Fine Arts, Paris. [Q to A 1996? We will need permission from the owner of the portrait to use this and a jpeg, tiff, or eps image of at least 300 dpi].

\textbf{Context setting and training}

Another informal contribution to research made by people with lived experience of dementia is enabling others to understand more of the context, place, and manner in which they live, or support someone to live, with their condition. Sometimes this takes the form of permitting researchers to observe the person in different scenarios, be it at home, while participating in an arts activity, or during clinical assessment. On other occasions the sharing may be more direct—through testimonies, giving talks, participating in demonstrations, or making audio and video recordings.

During the Created Out of Mind dementia residency, team members benefited greatly from the opportunity to observe dementia and arts practices. Seeing the effects of improvisatory music on people with advanced dementias participating in the Music for Life programme run by Wigmore Hall directly shaped the team’s research priorities and objectives (fig 2). As a result 360° camera based observational and quantitative rating techniques were developed to capture aspects of social interaction and musical, emotional, and physical engagement.\textsuperscript{18}
Project design and development

Shaping research design and development by participating in focus groups and advisory panels is an established component of public involvement. In recent years, co-production, referring to service providers and users working together to reach a collective outcome, has increased, particularly in translational research. In some cases, even formal involvement in co-design is founded on a much broader set of experiences shared between the person with lived experience and the researcher. Testimonies of researchers suggest that important ideas shaping research often emerge when people with a lived experience naturally and informally adopt an investigative or reflective stance in response to the challenges they face, which sometimes involves applying their previous professional skills to a problem area.

Two proponents of this kind of involvement in the co-design process are Pam and Richard Southerden, who have brought their experience of working in the IT industry to bear on technologies to support people living with dementia since Pam’s diagnosis with PCA in July 2015. Working with neuropsychologist Aida Suarez Gonzalez and developer Ashley Peacock, they have influenced the design of reading app ReadClear. They have also collaborated with computer scientist Nick Firth and social scientist Emma Harding to explore the potential utility of Amazon Echo voice controlled devices in the homes of people with dementia related visual impairment.

Extra benefits from formal participation

Involvement is typically distinguished from the many contributions people with dementia and their supporters make through participation (e.g., enrolling in research, providing supportive information relating to a participant, supporting the participation of another person by accompanying them on a research visit). However, the benefits of participation often exceed the scientific objectives of the study.

In some cases, participation leads, intentionally or unintentionally, to contributions to research which go beyond the data acquired. For example, during a routine verbal memory test conducted as part of a drug trial, one participant with PCA made a surprising observation. Presented with the first in a series of printed words to read and remember—SAND—she commented, “I can see an ‘S’ and I can see a ‘D’ but I can’t see the letters in the middle.” Her sharing of this hitherto unconsidered difficulty led to the design of experimental tests of spaced and unspaced letter identification (fig 3), the discovery of excessive visual crowding.
(difficulty identifying objects in clutter) in PCA, and the design of reading aids for people with dementia related visual impairment.

**Fig 3** Flanked letter identification tests for visual crowding. Participants are asked to identify a central target letter surrounded by flanking letters, symbols or numbers (adapted from Yong et al). People with excessive visual crowding can perceive the target letter reliably only when the flankers are spaced out (bottom line)

Just as formal involvement has been found to benefit both researchers and volunteers, so informal contributions can be valuable to all parties concerned. After reading a draft of the current article, Simon Rosser, whose wife had PCA, put into words what she felt: “The things that you are talking about, patient participation in research, patient groups such as the support group, particularly where the condition is not capable of being cured or treated in any meaningful way, is that the patient feels they are making a contribution and that they are doing something. Turning a natural feeling of helplessness in these situations into one where the patient is helping and feels that they are doing so gives the patient an enormous psychological boost. The very fact that people are interested in their views and in them (not just as a doctor treating them) and are wanting to understand them and what they are going through means a great deal to them.”

**Creating opportunities**

The examples of informal research contributions here all relate to the dementias but have equivalents in many other areas of healthcare. All the examples have had a big effect on the research programme, but they are serendipitous and not predictable. Various strategies can increase opportunity for informal input into research (box 2), but these should be regarded as complementary not alternative to more established formal involvement and the structures, processes, and definitions that support it.

**Box 2: Practical suggestions for gaining informal input**

- Maximise opportunities to meet people living with a health condition—eg, sit in on a clinic where people are describing their situation or attend care homes open days and support groups. Basic scientists may find taking part in local science festivals fruitful
- Learn about lived experiences online. There are many patient and carer forums, often run by the disease charities, where people share their experiences, coping strategies, and solutions that provide a trove of potential informal inputs
- Ask open questions when meeting people—eg, “What is it you don’t understand about your condition?” and follow the conversation from there
- Be open to listening to all aspects of experience. Expect the unexpected, not just those symptoms, experiences, and questions traditionally recognised as part of the condition

Collectively the stories of informal as well as formal involvement constitute an appeal to enable people affected by health conditions to influence the research which ultimately may affect them. In a world where research can often feel sanitised, these stories also encourage
researchers to recognise how subtle interactions can influence emerging ideas. As Arends and Thackara wrote:

“Science is hypothesis driven … but the first step on this journey—namely, the generation of the hypothesis itself, is rarely discussed… This early and ill-described stage entails a ‘methodology gap’. It is at this messy and amorphous juncture, where the creation of knowledge starts, that scientists must find their inspiration.”

In our experience, that inspiration often emerges through the experiences, questions, and uncertainties of those with a lived experience of a health condition.

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