SPOTLIGHT: PATIENT CENTRED CARE

Time to deliver patient centred care

We must harness the energy, insight and expertise of patients, carers, and the communities that support them to help drive change

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Patient centred care is central to the mission of healthcare, yet traditionally neither patients nor the public have had the power to shape the services they use and pay for, or define their value. As a result, many patients find services difficult to navigate, disempowering, burdensome, and seemingly designed to frustrate.

There have been repeated calls to enlist patients’ help to reform systems poorly geared to meet current health challenges, subject to political meddling, and permeated by competing interests that fuel overdiagnosis and overtreatment. UK and US policy supports person centred care and shared decision making, and the World Health Organization champions it. We know it supports person centred care and shared decision making, that fuel overdiagnosis and overtreatment.

There is wide support for measuring and incentivising person centred care. In New South Wales districts are competing to provide it, with promising results. Robert et al describe how to code design services with patients and carers. And the familiar message that strong leaders who ensure that all staff are responsible for patient partnership and person centred care can transform failing services fast is repeated in a podcast.

It’s agreed that feedback from patients and their families should be more rigorous and used to inform practice, not merely collated for research. Nelson illustrates the value of systematically collecting information on patients concerns and priorities at the point of care and using outcome measures that they identify.

Care should promote self management, not dependency

The rising number of people living with long term conditions, multimorbidity, and frailty requires services to be radically redesigned, and Eaton and colleagues highlight new approaches. These have to be based on a better understanding of what people need from health and social services, and patients can be crystal clear about this.

People who live with long term conditions are already in charge of their own health and if given the chance, willing to take on even complex management tasks. But it won’t happen without the development of the science of care and support planning, better training of front line clinical staff, and inclusion of patients and carers in multidisciplinary teams. Tailored support reduces dependency and can help the disadvantaged most.

All services should keep the burden of treatment on patients as low and “minimally disruptive” as possible.

We also need to find better ways to listen to patients and ensure their voice is included and heard in medical forums. Better conversations are also core to realising shared decision making based on individual priorities and preferences. The quality of the information on which joint decisions are based needs to be improved too, and innovative decision aids hold promise.

Digital revolution is changing traditional roles

The democratisation of health information, use of social media, rapid growth of networked patient communities, and new technologies have changed the landscape and provide new opportunities to harness the energy and expertise of patients. Clinicians and medical educationalists must catch up.

Empowered, engaged e-patients are growing a social movement and spearheading a shift in roles “as profound as women’s
liberation,” says de Bronkart, who suggests we need a new science to understand and optimise the role of the patient.27 Although the global flow of information has transformed many aspects of our lives, it has bypassed chunks of the health sector and it is still hard to get full access to personal health records. But there is welcome news on how the international move to open notes to patients is gaining momentum.28 However, it needs to go beyond giving people “viewing rights.” Montori emphasises that indigestible records must be made more useful to patients (and doctors) and foster communication, not substitute for it.29

In an unequal world the issue for billions of people is not whether care is patient centred but whether it is available at all, its poor quality, and its high cost. Anger and disillusionment are spurring change, however, and Jain provides illuminating examples of how patients’ rights are being advanced by effective advocacy in India, and local communities are holding the health sector to account.30 It’s time to get real about delivering person centred care. It’s not a panacea for all of medicine’s ills, but we should not underestimate its contribution to tackling them. Working collaboratively and sharing decisions about care, services, and research is challenging. It requires a sea change in mindset among health professionals and patients alike. But its rewards are rich and reaped mutually.

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