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What is the point of citizen participation in health care?

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Abstract

Does citizen participation in health care planning and provision have a clear purpose? Can it reasonably be considered a unified phenomenon? Current conceptual accounts – including a range of typologies descended from Arnstein’s ubiquitous but misunderstood ‘ladder of participation’ – are inadequate. The popularity of citizen participation belies fundamental uncertainties about what it entails and its associated benefits. A more pragmatic and less normative usage of the term is vital for the future.

Participation has been a central if indistinct part of our understanding of good health care since at least the WHO's Alma Ata declaration, which stated “The people have a right and duty to participate individually and collectively in the planning and implementation of their health care”.\(^1\) Since then an industry of commentators and practitioners has emerged. However, disagreements about the basics of participation remain unresolved and frequently unaddressed. In current practice, staffing public education stalls, attending presentations on car parking and filling in surveys on patient experience are seen as participatory acts,\(^2\) yet the rhetoric of participation remains rooted in power and conflict. Does participation in health care have a clear purpose and can it be considered a unified phenomenon?

Introducing participation

While the concept of participation is widely recognised as difficult,\(^{4-6}\) literature on participation in health care has continued to grow apace. One result of this is a remarkable
degree of terminological instability. Debate revolves around a range of terms formed by adding a named group of participants (‘public’ ‘patient’ ‘citizen’ ‘community’ or ‘user’) to a type of activity (‘involvement’ ‘engagement’ or ‘participation’). This is not merely semantic but denotes disagreement about the nature and purposes of participation.

Attempts to define participation in health care end up unworkably vague. For example, Titter’s effort to conceptualise ‘public and patient involvement’ (the preferred term in the English NHS since the late 1990s), ranges from direct public decision-making to the gathering of patients’ views on everything from individual treatment decisions to the training of health professionals. It seems that almost any health care activity involving a conscious patient or citizen can be understood through the lens of participation. Titter seeks to contrast public and patient involvement with patient choice but even this is not obviously excluded from the definition he offers: “Ways in which patients can draw on their experience and members of the public can apply their priorities to the evaluation, development, organization and delivery of health services”.

Titter is not alone in struggling to produce a workable definition of participation. Abandoning the search for a catch-all description, numerous authors have turned instead to a system of classifications to deal with the vast range of initiatives that profess to be ‘participation’. These typologies emphasise the degree of power gained by citizens, the public, or specific service users. Existing typologies can be mapped against each other and, even where the label used differs, the content of each ‘level’ of participation is reasonably consistent. However, in a demonstration of the risks of a literature emerging without a solid conceptual grounding, over time typological levels have tended not just to reduce in number but to concentrate in the middle of the ladder. Arnstein’s original ‘ladder of participation’ ran
from citizen control to the bottom rungs of manipulation and ‘therapy’ (an interesting
inclusion given the trend for participatory initiatives to be sold as producing more informed
and responsible service users). More recent literature ignores these provocative possibilities
and presents instead a technical ‘menu’ of approaches. In this way, typologies of participation
have become less radical over time and have diluted Arnstein’s highly critical account.

Why should participation be a priority for health systems?
In health care systems, potential (claimed) benefits for citizen participation range from
accountability for public funds, medical rationales for responsiveness (incorporating the
plethora of potential gains from a ‘co-production’ approach), consumeristic objectives of
service efficiency, better informed (and implicitly more ‘responsible’) service users, and
mutual support for service users. Bochel and colleagues’ review of the benefits of
participation argued that: “participation might be intended to improve governance,
democracy, social capital, education and development of individuals, policies, service
implementation and delivery – all, or one or more of these, or something else altogether” 11.
In short, participation is presented as offering better services, better outcomes and better
service users. The authors nonetheless conclude by re-stating their commitment to the
possibility of “real” participation: like Tritter 6, they evidently operate with a tacit
understanding of the purposes of participation and yet a clear statement of these is not
offered.

Harrison and colleagues10 depict three versions of participation in health care: consumerist,
democratic and ‘New Social Movement’ (an emancipatory concern with vulnerable or
marginalised sections of society). These can be understood as groupings which reflect deeper
divisions in approach and purpose. The concerns of (consumeristic) public management
scholars are often rooted in organisational effectiveness, which is assumed to be a broadly straightforward (and objectively determinable) goal: as long as someone is voicing concerns about an organisation, it is not of great concern whether they are representative of the wider population.

Researchers working from the perspective of New Social Movements would likely dispute this. For these researchers, the lessons of the disability movement and other user groups are that mainstream preferences for an organisation can be profoundly at odds with what some of the most vulnerable sections of the population need. Arnstein’s arguments, built from a concern with society’s ‘have nots’, fall into this category and, when next we cite her ladder of participation, we should do so with awareness of the particular radical perspective which has made this work so widely known and so poorly understood. Finally, for scholars concerned with an overarching picture of democracy, rather than a primary emphasis on vulnerable groups, concerns are about “the fundamental democratic tests of popular control and political equality”.

While good analysis may produce a clearer specification of participation, this would require academics, policy-makers and practitioners to agree on what they understand as participation in health care. Alternatively, participation can be seen as “a descriptive umbrella term for the spectrum of processes and activities that bring the public into the decision-making process”. While this might sacrifice the aura of righteousness which currently surrounds participation, there are advantages to acknowledging the range of participatory activities and rationales, accepting differences and being clear about them.

**Conclusion**
The persistent yet ill-specified normative appeal of participation in health care is undermined by a lack of conceptual foundation. The lack of specification of participation allows policy-makers, practitioners and researchers to go about their business without having to resolve the uncertainties and conflicts contained within the phenomenon, or to choose between the democratic, consumeristic or emancipatory rationales discussed above.

Abandoning the idea that participation in health care denotes a self-evident (and self-evidently commendable) set of activities requires us to be much clearer about the purposes of participatory activities and the standards by which we judge them. Doing so will force us to acknowledge the changes in emphasis which can be identified over time. Incoherent definitions have facilitated the growth of both literature and policy on public participation in health care. It is time to admit that participation rationales are far from self-evident and accept that participation is a site of disagreement, rather than an unqualified good.


