On behalf of the authors, the reviewers, the editorial staff and our wonderful Editorial and Advisory Boards I extend a warm welcome to our first issue of *Qualitative Research in Medicine and Healthcare* (QRMH)! I am lucky to be working with so many great people, and dedicated to making this a great place to publish.

With many qualitative research journals already on the academic scene, and a good amount of these in the areas of health communication, healthcare, and medicine a new journal is a risky proposition. As our aims and scope promise, it is nonetheless a vital and crucial undertaking. As the articles in this first issue demonstrate, QRMH is true to its promise of interdisciplinarity; four of its five pieces are collaborations by scholars and practitioners in medicine, nursing, communication, biology, theology, international affairs and conflict resolution. And all of the articles raise complex queries, tell difficult stories or ask the complex and uncomfortable questions for which qualitative research is best equipped.

As many of you already know, qualitative research is not for the faint of heart. It is reflexive, positioned and like life itself, often messy and focused not on producing easy answers but on the very process of questioning. It is, at best, acutely aware its ontological consequentiality. Unlike research studies under positivist or post-positivist auspices, true qualitative research does not separate the observer from the lived experience of those whom she observes.

Rather, it is by understanding the ways in which our own research practices bring forth the very world in which we live that qualitative researchers have a stake in understanding the very reflexive dynamics of how we constitute the world makes us.

Consider the traditional process of knowledge construction in medicine and healthcare and how, in these very social and human sciences, we unproblematically take notions of scientific evidence as detached from the very measurements, screenings, and processes of lived experience in which knowledge-making is situated. Data, findings, reports are always written in the third person omniscient language of *science*, concealing writers, bodies, lives. We try to *fit* ourselves into the spaces offered by questionnaires, rather than finding questionnaires designed to fit us.

Though *evidence-based* has become a popular phrase, for example, it works by pronominal omission; and yet it is always evidence for someone, by someone, and in a particular context with very real and life changing consequences. By the same token, that of *patient* (including the popular *patient-centered*), *disadvantage*, *vulnerable*, and so on are constructs – that is, categories that create the very situations they purport to describe. By way of our research *findings* and discussions, these categories can either re-enter the world as natural, or we may use research as a process in which to examine the social dynamics of what it means for these terms to categorize and create what Wittgenstein called *forms of life*, that is, to describe, as well as prescribe conditions of our very being and courses of action that implicate courses of actions, policies, social change.

As Erin Castelloe tells us in an intensely personal and beautifully written opening article (and the second part to this first one is already in the works!) she is a woman at a crossroads, facing important decisions as to how to (re)construct her career and personal life. Metaphorically, working in healthcare is a matter of always being at a crossroads. Medicine, she writes, is practiced by imperfect human beings, most of whom are dangerously overworked. The stories that enrage me are those in which doctors dismiss people’s concerns without a complete assessment: the man without insurance who is obviously having a heart attack but is discharged from urgent care.
The young woman with new-onset, nocturnal seizures who is diagnosed with a supra-tentorial issue (anxiety and/or personality disorder) without a neurological exam, let alone a sleep study and nocturnal electroencephalogram (EEG).  

The study sheds new light in the way policy makers may employ accounts to make effective changes in communities where health insurance is not used as proponents wish it could be.

The final article in this issue, by Gathercoal, Gathercoal, Seegobin, and Hadley combines quantitative and qualitative approaches in an innovative and creative research design. The authors challenge the view that disabilities are either romantic conditions that render those affected with them special in some way or, on the other hand, horrible biological or physical handicaps that dehumanize them instead of seeing these individuals as ordinary people facing extraordinary, and often society-created obstacles. I find no better way to conclude this introduction than to use these authors commentary on reflexivity, constitutiveness, and consequentiality in research, and the fearlessness of qualitative researchers to be aware of these issues. As Gathercoal et al. so aptly note,

Even traditional concepts such as effective come embedded with specific cultural and social perspectives with biases, adopting positions which take for granted ill-defined social goals and unintended consequences. Who, for example, gets to determine just what is best for both clinicians and for clients?

In leaving you to this and many more thought provoking questions and insightful studies that compose this first issue, I extend an invitation to all of you who are reading that you will submit to our journal, and join in the exciting, innovative, rigorous and spirited research conversations that will take place in this forum.

References