Chapter 7.3
The Impact of Privacy Legislation on Patient Care

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ABSTRACT
This article looks at how privacy legislation in Canada may interfere with patient care and the use of electronic health records. A review of the literature shows that legislation across Canada is not uniform and varies to the degree in which it addresses issues of privacy and confidentiality. What is not clear is what impact legislation has on the movement towards electronic health records? A case study was undertaken to look at a specific project involving the design of an electronic health record as a means of sharing information between physicians and prostate cancer patients. Through interviews with those associated with the project, it was clear that legislation itself was not a barrier. The problem was that the legislation was open to interpretation. The author hopes that an understanding of the issues raised by this article will lead to further discussion and research on this topic.

INTRODUCTION
Patient privacy is a topic of great interest both in Canada and internationally. Many health conferences and academic journals often have several papers devoted to this topic. This issue has gained more attention in the last five years as there is a continuing trend in health care towards more electronic capture and sharing of information between providers and patients. It has been recognized that health care has typically been behind other industries, such as, airlines and banking, when it comes to embracing its users in the sharing of electronic information. Patients and their health care providers have expressed an interest in sharing of information to facilitate optimal patient care. Unfortunately, as more patient information is made available electronically, there is an increased potential for loss of privacy and confidentiality.
In Canada most provinces have legislation to ensure that there are safeguards in place to protect confidentiality and security of sensitive personal data. However, it may be that this legislation is too restrictive and interferes with patient care activities. Many initiatives, such as those led by federal and provincial governments, are looking at novel ways of providing access to and sharing of electronic patient information. Often these initiatives involve the use of the Internet or some form of information and communication technology (ICT). These initiatives may run into issues with current legislation, to the extent that they never get off the ground or at best are significantly delayed in implementation. In general, the topics presented usually deal with how to be compliant with legislation or how to go about addressing issues related to privacy. Little has been discussed that addresses how legislation may have an impact on patient care. Indeed, there is scarce information in the literature that looks at how the legislation in any particular jurisdiction may have an impact on novel approaches to the delivery of care.

The aim of this article is to look at a new and innovative project for the sharing of information for prostate cancer patients in the Canadian province of British Columbia (BC). A case study was undertaken in order to examine whether current legislation has had an impact on this project, known as the Provider Project.

BACKGROUND

The Provider Project

The British Columbia Cancer Agency (BCCA) provides cancer care programs for the people of BC, including prevention screening, early detection, diagnosis and treatment services, support programs, community programs, research, and education. BCCA, along with seven other provincially mandated health care agencies, is a member of the Provincial Health Services Author-
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