Handbook on Sensitive Practice for Health Care Practitioners:

Lessons from Adult Survivors of Childhood Sexual Abuse
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This may be a person who’s gone through something very traumatic ... [who needs] some really safe technique ... Because otherwise you’re going to have a certain segment of patients that are going to walk away feeling as though they’ve been abused all over again, quietly abused, just walking away and seeking another health care practitioner, just going through the cycle, again and again and again, and maybe not understanding why, maybe not knowing how to say it, how to voice that, just keep going through that whole cycle over and over again. There’s a huge populace out there that just needs that extra gentle care. It’s because of that, maybe the whole populace needs to be treated the same way.

— A male survivor of childhood sexual abuse —
1.1 Audience and focus

This handbook presents information that will help health care practitioners practise in a manner that is sensitive to the needs of adult survivors of childhood sexual abuse and other types of interpersonal violence. It is intended for health care practitioners and students of all health disciplines who have no specialized training in mental health, psychiatry, or psychotherapy and have limited experience working with adult survivors of childhood sexual abuse. This second edition includes experiences and ideas of both women and men survivors as well as of practitioners from more than ten health disciplines. The Handbook is not meant to encourage health care providers to step outside their scope of practice, nor is it a substitute for the specialized training required to provide intensive psychotherapy or counselling for survivors.

The Handbook is based on extensive interviews, group discussions, and a national consultation process involving adult survivors, clinicians, and mental health practitioners. Direct quotes from participants are included to illustrate selected issues and to connect health care providers in a more personal way to survivors’ thoughts and feelings. To our knowledge, the Handbook is the only work in print that has employed a process of bringing adult survivors and health care practitioners together to develop an empirically grounded account of the issues and problems that adult survivors of childhood sexual abuse encounter in health care settings. The empirical basis of the Handbook is found in Appendix A.

1.2 An issue for all health care practitioners

As many as one third of women and 14% of men are survivors of childhood sexual abuse. Childhood adversity – including sexual, physical, and emotional abuse – is associated with a greater risk of a wide variety of health problems. This means that all health care practitioners – whether they know it or not – encounter adult survivors of interpersonal violence in their practices. Survivors are health care consumers of every age who seek all types of health services, and our hope is that the principles and guidelines of Sensitive Practice will become “universal/routine procedures” in all health care encounters and that all health care consumers will benefit from them.

Section 2.5 – Childhood sexual abuse and health

Examinations and procedures that health care providers might consider innocuous or routine can be distressing for survivors, because they may be reminiscent of the original trauma. Exclusive focus on the body, lack of control, invasion of personal boundaries, exposure, vulnerability, pain, and sense of powerlessness are common experiences in the health care environment and may be extremely difficult for survivors because they can mirror aspects of past abuse. An appreciation of the dynamics and long-term effects of childhood abuse is the first step toward a better understanding of survivors’ needs and responses to care. Sensitive Practice builds on core competencies to help health care practitioners be more understanding of and responsive to the
specific needs of adult survivors of violence and abuse.

Chapter 2 – Background Information about Childhood Sexual Abuse
Chapter 3 – What Childhood Sexual Abuse Survivors Bring to Health Care Encounters

Although our research focuses primarily on Sensitive Practice in traditional health care settings, health care providers work in widely diverse areas including client homes, rural and remote areas, and school systems. We believe the principles and guidelines of Sensitive Practice can be adapted to all health care venues.

Chapter 3 – What Childhood Sexual Abuse Survivors Bring to Health Care Encounters

1.3 Organization

The Handbook is divided into nine chapters. Readers are alerted (watch for a blue “i” in a blue circle) when the topic addressed in one section is clarified or expanded upon in another section.

- Chapter 1 offers an introduction to terminology and suggestions for using the Handbook.
- Chapter 2 provides basic information about the nature and scope of childhood sexual abuse and health problems associated with a history of childhood sexual abuse to assist health care practitioners to understand the significance of Sensitive Practice.
- Chapter 3 presents information about how experiences associated with a history of childhood sexual abuse may be manifested in health care settings.
- Chapter 4 describes the principles of Sensitive Practice derived from our research; we consider these principles foundational to ethical health care.
- Chapters 5, 6, 7, and 8 outline the guidelines for Sensitive Practice. These guidelines operationalize the principles of Sensitive Practice and are meant as practical suggestions that health care practitioners can incorporate into their clinical practice.
- Chapter 5 presents those guidelines related specifically to the context of health care encounters.
- Chapter 6 details guidelines applicable to all interactions between health practitioners and their clients.
- Chapter 7 offers health care practitioners guidelines for coping with the problems that can occur in encounters with child sexual abuse survivors.
- Chapter 8 highlights guidelines focused on disclosure of past abuse, especially as it is related to survivor-clinician interactions.
- Chapter 9 explores the contributions which health care practitioners can make to an adult survivor’s healing and recovery, and elaborates on the relationship between Sensitive Practice and patient-centred care.
- Chapters 4, 6, and 8 conclude with questions intended to stimulate reflection about the application of the principles and guidelines to health care practices.
- Nine appendices augmenting the text and a bibliography detailing both works cited and recommended readings and resources finish off the Handbook.

1.4 Suggested uses of this Handbook

Students, practitioners, and administrators are urged to think about the information on violence and abuse and Sensitive Practice provided in this Handbook, and to reflect on its potential...
for informing their own practice and workplace policies. Specifically, health care providers should consider:

- How the information applies to them;
- How to implement the principles and guidelines into their practices;
- How they might best respond to the various situations described in the Handbook.

We believe the information in the *Handbook* applies to everyone in health care environments. Many of the difficulties that adult survivors experience in these environments arise because practitioners who work in them are unaware of the effects of violence on health and health care or because organizational policies and attitudes have not taken this information into consideration. Before dismissing a suggestion as inapplicable to their practices, clinicians are encouraged to reflect upon the following questions:

- What aspects of this suggestion do not apply?
- If a suggestion is not completely applicable, what element(s) of it could be relevant?

The Handbook can be used to help health care practitioners:

- Learn more about the effects of interpersonal violence on health;
- Work more effectively and compassionately with affected individuals;
- Identify and respond sensitively to individuals who are triggered or dissociate in a health care encounter;
- Feel better prepared to work with patients who disclose past abuse;
- Teach administrative personnel and assistants about childhood abuse and its implications for their work;
- Disseminate ideas from the Handbook to colleagues with the intent of creating an integrated and responsive network of care;
- Influence policies and practices within public agencies to be more sensitive to survivors;
- Coach students and colleagues to critically analyze professional practices;
- Reflect on their philosophies of care and how they are expressed in day-to-day practice, with the intent of becoming more ethical, congruent, and sensitive in their work;
- Develop clear guidelines to address their concerns about best practice with a specific patient or treatment procedure.

### 1.5 Terminology

The following is a clarification of terms the reader will encounter in this *Handbook*. Many health care providers have various preferences for the words *patient* and *client, clinician*, and *practitioner*, and for the terms they use to describe their work (examination, treatment, etc.). However, all readers are encouraged to look beyond the terminology to fully consider how the information applies to their own practices. 

*Survivor* or *adult survivor* is used instead of *victim* when referring to adults who have experienced childhood sexual abuse to acknowledge the strength and resourcefulness of individuals who have lived through the experience.\(^{23}\) Attitudes about the words *survivor* and *victim* vary among those who have experienced childhood abuse, as well as among those who work with these individuals.
range from eating well and exercising regularly to adhering to clinicians’ specific recommendations.

- Participants’ words appear in *italics*.

### 1.6 Limitations

The experience and long-term effects of childhood sexual abuse are affected by a complex interaction of factors including: (a) those related to the individual (e.g., genetics, stage of development at which the abuse occurred, personal coping resources); (b) the abuse itself (e.g., frequency, duration, relationship between perpetrator and victim); (c) the presence and quality of social support at the time of the abuse and into adulthood; and (d) those related to the larger environment, including culture, ethnicity, and other social determinants of health. Adult survivors who participated in our studies were recruited from agencies, groups, and individuals offering counselling and support. Thus, they are individuals who have worked or are working towards recovery with the assistance of external support.

Notwithstanding the diversity and uniqueness of these participants, the *Handbook* cannot claim to address Sensitive Practice for adult survivors with every abuse experience, of every ethnicity and culture, of every sexual orientation, or at every stage of recovery. Similarly, although we have tried to address a wide range of health practitioners working in various settings by incorporating a broad consultation in the research method, we cannot claim to address every aspect of Sensitive Practice for every type of health care practitioner.

While acknowledging these limitations, we believe that this second edition of the *Handbook* presents a framework for working with adult survivors of interpersonal violence in all types of practice that is both accessible and empirically derived. We hope that health care practitioners will adapt and refine the guidelines as they work with survivors whose unique needs and reactions were not represented by the research participants.
2 Background Information about Childhood Sexual Abuse

2.1 Definitions

While the sexual exploitation of children and adolescents is a criminal act, legal definitions of childhood sexual abuse vary across jurisdictions. There is, however, wide agreement that childhood sexual abuse involves: (a) sexual acts with children and youth who lack the maturity and emotional and cognitive development to understand or to consent; and (b) “an ‘abusive condition’ such as coercion or a large age gap between participants, indicating lack of consensuality.” 26p.32 In general, children and younger adolescents are unable to consent to sexual acts with adults because of their lack of maturity and relative lack of power. An abusive condition implies a difference in power between the perpetrator and the victim. Children can also be abused by other children or adolescents who have more power by virtue of age, physical strength, life experience, intelligence, authority, or social location. The Canadian Incidence Study of Reported Child Abuse and Neglect tracked eight forms of child sexual abuse: penetration (penile, digital or object penetration of vagina or anus), attempted penetration, oral sex, fondling of the genitals, adult exposure of genitals to child, sexual exploitation (e.g., involving child in prostitution or pornography), sex talk (including proposition of a sexual nature and exposing a child to pornographic material), and voyeurism. 168p.38-39

An extreme and controversial type of abuse is ritual abuse, which has been defined as psychological, sexual, and/or physical assault on an unwilling human victim, committed by one or more individuals, as part of a prescribed ritual that achieves a specific goal or satisfies the perceived needs of their deity. 27,140

2.2 Childhood sexual abuse survivors

The great paradox of childhood sexual abuse is that, while it has become more prominent in the public consciousness, it remains shrouded in secrecy. Media coverage of high-profile disclosures and investigations provide evidence that childhood sexual abuse does exist – in “good” families and “trusted” institutions, at all socioeconomic levels, and among all racial and ethnic groups. Frequently we hear and read stories about survivors who are men and women from all walks of life – students, sports figures, clergy, entertainers, educators, police officers, judges, politicians, and health care practitioners. They are our friends and neighbours, our colleagues, and sometimes even ourselves or members of our own families. Despite this prevalence, most childhood sexual abuse survivors are invisible to us, particularly given that it is estimated that fewer than half disclose their abuse to anyone. 62,105 Some are silent because they fear reprisal from their abusers; others worry they will not be believed or that they will be blamed or even punished. 56,113 Still others say nothing because they harbour the erroneous belief that they are responsible for their abuse.

* According to the Criminal Code of Canada, when sexual activity is exploitive (such as sexual activity involving prostitution, pornography, or a relationship of trust, authority or dependency) the age of consent is 18. For sexual activity which is not exploitive, the age of consent is 16 years. The exceptions are that a 12 or 13 year old can consent to engage in non-exploitative sexual activity with another person who is less than 2 years older; and a 14 or 15 year old can consent to engage in non-exploitative sexual activity with another person who is less than 5 years older. A 14 or 15 year old can also consent to engage in sexual activity with a person to whom they are married. These laws governing the age of consent for non-exploitative sexual activity came into force on May 1, 2008. Transitional provisions allow 14 and 15 years old who were in common-law relationships on May 1, 2008, to continue engaging in non-exploitative sexual activity.
little consideration for their effect on the child. Some child abusers use physical force or explicit threats of harm to coerce their young victims into compliance, while others develop long-term relationships with their victims and carefully groom them with special attention or gifts. While childhood sexual abuse does not always involve physical injury, it is a violation of body, boundaries, and trust and is typically experienced as traumatic.

While people who report a history of childhood sexual abuse are at increased risk for a wide range of difficulties in adulthood, studies suggest that “in the region of 20% to 40% of those describing CSA [childhood sexual abuse] do not have measurable adult dysfunction that could be plausibly be related to abuse.” A number of factors affect how a particular individual may respond to childhood sexual abuse. Some of these include the gender of the perpetrator, the number of perpetrators, the nature and closeness of the relationship between victim and perpetrator, the duration and frequency of the abuse, characteristics of the abuse itself (e.g., contact vs. noncontact, penetration, etc.), the use of threats or force, and the age of the victim at the time of the abuse.

As many as one third of women and 14% of men are survivors of childhood sexual abuse.
Background Information about Childhood Sexual Abuse

Chapter 3 – What Childhood Sexual Abuse Survivors Bring to Health Care Encounters

Chapter 8 – Guidelines for Sensitive Practice: Disclosure

Table 1 lists the findings of a number of studies that have examined the correlation between histories of childhood sexual abuse and later health and function. Considerably more studies have examined these relationships in women, and when male survivors have been studied, the relationship between past abuse and the mental health of male survivors has been the primary focus. Guy Holmes, Liz Offen, and Glenn Waller argue that two pervasive myths – that males are rarely sexually abused and that childhood sexual abuse has little effect on males – deter boys and men from disclosing their abuse and, in turn, prevent society from legitimizing it as a problem.

Because most health care practitioners do not routinely inquire about childhood sexual abuse, its long-term effects are under recognized, its related health problems are misdiagnosed, and it is not met with a sensitive, integrated treatment response.

Two pervasive myths - that males are rarely sexually abused and that childhood sexual abuse has little effect on males - deter boys and men from disclosing their abuse and, in turn, prevent society from legitimizing it as a problem. The increasing societal recognition of the prevalence and seriousness of sexual abuse of boys is likely to lead to further investigation of physical health correlates.
### TABLE 1
Correlates of childhood sexual abuse and measures of health and function:
A selected list of findings from research studies

#### In females, a history of childhood sexual abuse or a range of childhood traumas including sexual abuse is correlated with:

- poorer physical and mental health and a lower health-related quality of life than non-traumatized individuals\(^59,145,176\)
- chronic pelvic pain\(^129\)
- gastrointestinal disorders\(^53,141\)
- intractable low back pain\(^146\)
- chronic headache\(^58\)
- greater functional disability, more physical symptoms, more physician-coded diagnoses, and more health risk behaviours, including driving while intoxicated, unsafe sex, and obesity\(^176\)
- ischemic heart disease, cancer, chronic lung disease, skeletal fractures, and liver disease\(^59\)
- high levels of dental fear\(^177,185\)
- greater use of medical services\(^35,87,102,116,150,178\)
- drug and alcohol use, self-mutilation, suicide, and disordered eating\(^56\)
- adult onset of 14 mood, anxiety, and substance use disorders\(^108\)
- higher rates of childhood mental disorders, personality disorders, anxiety disorders, and major affective disorders, but not schizophrenia\(^154\)
- diagnosis of Borderline Personality Disorder\(^79,82,100\)

#### In males, a history of childhood sexual abuse is correlated with:

- anxiety, low self-esteem, guilt and shame, depression, post-traumatic stress disorder, withdrawal and isolation, flashbacks, dissociative identity disorder, emotional numbing, anger and aggressiveness, hypervigilance, passivity and an anxious need to please others\(^22,28,32,45,147\)
- adult onset of five mood, anxiety, and substance use disorders\(^108\)
- substance abuse, self-injury, suicide, depression, rage, strained relationships, problems with self-concept and identity, and a discomfort with sex\(^49,54,133,161\)
- increased risk of HIV\(^5\)
- anxiety and confusion about sexual identity and sexual orientation\(^85,133\)
- increased risk of “acting out” aggressively\(^85\)
- contact with criminal justice system\(^85,96\)
A female abuse survivor may also be mistrustful of authority figures, which stems from having been betrayed by the trusted adult who abused her. This helps to explain the difficulty that some survivors have trusting health care practitioners and why they experience health care encounters as distressing. It also helps explain why so many female survivors report symptoms of depression and anxiety:

*I didn’t want to state what my needs were because ... [with] the abuse ... you don’t get to choose what happens to you. What happens to you happens to you, you just accept it and that’s the way I thought for a long time. I still probably think that way but I’m trying to change the way I think because I do have choice now.* (Woman survivor)

Girls learn that it is important for females to be objects of male sexual desire and that appearing young and innocent is sexually appealing. “We dress fashion models up to look child-like and sexually provocative and set this standard for all women,” writes Calgary social worker Lois Sapsford. Girls may also learn that, to be valued, they must be sexually “pure”; at the same time, they receive the contradictory message that they should be not only beautiful but also “sexy.” Sexual abuse objectifies a girl’s body to serve the needs of her abuser and may leave her believing that her sole value is as a sexual object. The message that females should be sexually “pure” along with the stigma attached to sexual abuse contributes to some female survivors’ perceptions of themselves as “damaged goods” and to the shame and guilt that many describe. This may be manifested in a survivor’s ambivalence about her body and reticence to seek care for health problems:

*Most survivors I know hate their body, disown their body ... become disconnected from it.* (Woman survivor)
The other thing is the big shame and the secret ... We may have an ailment that could be addressed ... [early] but let it go and let it go until ... it takes longer to mend or to heal.

(Woman survivor)

The historical and current societal factors that encourage people in our society to deny or minimize the significance of child sexual abuse also affect female survivors’ perceptions about the wisdom of disclosing their experience. Many women participants talked about their fear of not being believed; some gave examples of being told directly that they must be lying or imagining things. Another aspect of female gender socialization is the message that it is the female who is responsible for setting limits on sexual behaviour, which contributes to women survivors fearing that they will be blamed for what happened, even though the sexual behaviour occurred when they were children and the perpetrator was older and more powerful. One health care practitioner responded to a woman’s disclosure of past abuse by asking, “How did you let it happen?” These societal messages strongly discourage women survivors from sharing their experience with health care practitioners, which in turn impedes the clinician’s ability to assess all factors that may contribute to health problems.

3.2 Gender socialization: Men’s experiences

The men in our studies repeatedly reported feeling invisible as survivors of childhood sexual abuse. Among the major factors contributing to the invisibility of male childhood sexual abuse survivors are: (a) the widespread lack of knowledge about the prevalence of childhood sexual abuse of boys; (b) incongruence between society’s notions of masculinity and victimhood; and (c) the fact that services for childhood sexual abuse survivors, which grew out of the second wave of feminism, were historically designed for women and not for men.

For a man to acknowledge that he has been sexually abused is an admission of vulnerability in a society that has few models for the expression of masculine vulnerability. Indeed, applying the label victim of sexual abuse to a man juxtaposes vulnerability with masculinity, an uneasy pairing that further contributes to the under recognition and underreporting of childhood sexual abuse among boys and men.3,43,52,86,105

The socialization of men to be strong and independent15,85 complicates the situation for male survivors who consider sharing their history of abuse with a health care practitioner.165 As Michel Dorais puts it in his book Don’t Tell: The Sexual Abuse of Boys, the “masculine conception of virility is incompatible with the factual experience of having been a victim of sexual abuse, or needing help following such a trauma”52p.17 (see also O’Leary117). Men in our study spoke about their need to appear “tough” and “in control” despite feeling anxious and fearful during encounters with health care practitioners:

Men are tough. Men are macho. Men don’t need [help]. All we have to do is to “get over it! Get over it – be a man!” You know, men don’t cry. (Man survivor)167p.509

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The stigma attached to sexual abuse contributes to some female survivors’ perceptions of themselves as “damaged goods,” as well as their ambivalence about their bodies and reticence to seek care for health problems.

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Some participants also spoke about their difficulty in identifying and expressing their feelings:

Women appear to me more aware of the names of things. Such as “I’m feeling depressed” or “I’ve been having a real struggle for the past couple of weeks and these are the
There is a pervasive belief that boys and men are rarely victimized and that a central feature of masculinity is the ability to protect oneself (Mendel as cited in Lab, Feigenbaum, & De Silva167); failure to do so is seen as evidence of weakness and can be a source of great male shame. Thus, the “dissonance between the male role expectation and the experience of victimisation”117p.83 may seriously compromise the health care of male survivors, often because their feelings of shame and unworthiness affect their ability to seek care:

“One of the reasons why for a long time I didn’t go [to a health care practitioner was that] ... quite frankly, I just didn’t feel worthy ... Worthy of the care, the attention. I mean doctors are busy. (Man survivor)

Most of the men in our studies expressed the belief that different reactions to male and female childhood sexual abuse survivors shape their help-seeking behaviours and, in turn, influence how health care practitioners treat them. In general, the participants suggested that health care providers are sceptical about men who disclose sexual abuse and tend to take their experiences less seriously than those of their female counterparts. In addition, some regard sexual abuse by a woman as something that the “fortunate” male survivor should have enjoyed. Ramona Alaggia3 and Guy Holmes and colleagues85 reiterate that such perceptions are common. The media also contribute to these views by framing the sexual abuse of boys by adult women as a “sexual relationship” (e.g.,36,149). The fact that boys are more often sexually abused by a female than girls31 may fuel the myth that sex between boys and women is normative rather than abusive and perpetuates the “male gender role of seeking early sexual experiences with women.”15p.225

Notwithstanding the general progress made in addressing homophobia in our society, some of the men in our study talked about their fear that health care practitioners would think they were homosexual if they revealed their history of childhood sexual abuse. Others talked about how their abuse experiences had led them to develop strong negative feelings about individuals (including health care practitioners) whom they perceived to be homosexual:

“I had to go into the hospital where I had a problem with some medication I had [taken] and there was a male nurse there and he was obviously very effeminate, and he had to give me an IV, I refused him because I didn’t want him touching me. (Man survivor)167p.506

Such reactions can be seen as internalized homophobia. These fears may also reflect the pervasiveness of the myth in our society that childhood sexual abuse causes boys and girls to become gay or lesbian.132

3.3 Societal myths about the cycle of violence

The emotional cost of childhood victimization is intensified especially for male survivors by the societal belief that it is only a matter of time before they become abusers themselves, if they have not already done so. The media typically give more attention to the erroneous belief that male survivors will likely become perpetrators43 than to information that disputes this belief.117 Despite the lack of conclusive evidence regarding this causal link (e.g.,68,137) and the fact that many male perpetrators do not report a history of childhood sexual abuse,99 the public and even some male survivors themselves continue to fear that they are destined to become perpetrators.85,117,133 Some female survivors may also fear that they will sexually abuse children or that others will see them as potential offenders.
3.4 Transference and counter-transference

The concepts of transference and counter-transference were originally identified by Freud in the context of psychoanalysis, and refer to common human experiences that are important for everyone working in human service to understand. **Transference** is said to occur when an individual displaces thoughts, feelings, and/or beliefs about past situations onto a present experience. It is widely agreed that we all engage in transference to some extent. While transference can be positive or neutral, it can also be negative and may interfere with healthy and adaptive functioning. For example, an adult who was constantly criticized by an authority figure may grow up expecting all authority figures to be critical and may hear criticism where none is intended. Similarly, survivors of childhood sexual abuse may react negatively towards a health care practitioner whose appearance, gender, or mannerisms are reminiscent of someone who abused them.

The dynamics of transference help explain why a survivor may respond to an interaction with a health care practitioner in ways that are unrelated to the encounter or to the specific health care practitioner. Understanding transference may also help health care practitioners to avoid taking patients’ negative responses personally.

**Counter-transference** involves the same dynamics as transference, but occurs when a health care practitioner responds to a patient with thoughts, feelings, and/or beliefs associated with his or her own past. For example, a patient who reminds a practitioner of an angry and demanding teacher may evoke feelings of anxiety that seem out of proportion to the current situation.

Counter-transference can also refer to the health care practitioner’s expectable emotional reaction to a patient’s behaviour – in particular, when the patient is transferring experiences from the past. For example, a survivor may engage in transference by behaving in a hostile manner towards a practitioner whom he incorrectly believes does not care about him just as his parents did not seem to care about his well-being. A health care practitioner who responds with anger and defensiveness can be said to be allowing counter-transference feelings to be expressed.

While it is understandable that health care providers have negative feelings in response to a patient’s negative transference, they must strive to contain these feelings and respond professionally. Inquiring about the reasons for the patient’s hostility, for example, is likely to be more productive than responding with anger. Health care providers have an ethical obligation to work continuously at being self-aware and to reflect critically on their practice in order to recognize when they may be responding harmfully to a patient’s transference or experiencing counter-transference. Further, health care providers need to remind themselves repeatedly of their obligation to respond to a patient professionally, even when they believe they have been judged harshly, have been provoked, experience negative feelings about the patient or are personally upset. When practitioners have difficulty meeting these ethical requirements, they need to reflect on the situation and the reasons for their responses and take appropriate steps to prevent harming their patients directly or indirectly. If a health care practitioner notices a recurring strong reaction to a particular individual or to certain behaviours, personal characteristics, or events, it may be useful to talk to a supervisor or trusted colleague about it.

3.5 Specific behaviours and feelings arising during health care encounters

**Distrust of authority figures.** Throughout this project, survivors told us how, as children, they experienced violation at the hands of an authority figure and how the distrust from these
Past experiences affect their interactions with health care practitioners. Although this distrust originates in the past and should not be taken as a personal affront, survivors constantly scrutinize health care providers for evidence that they are taking active and ongoing steps to demonstrate their trustworthiness. It is crucial to recognize that some survivors may associate a health care practitioner’s attempts to verbally assure them that they are safe with the perpetrator’s empty assurance of safety during their abuse.

Section 4.1 – Overarching consideration: Fostering feelings of safety for the survivor

Fear and anxiety. Many survivors spoke at length about their tremendous fear and anxiety during health care encounters. The experiences of waiting, being in close contact with authority figures, and not knowing what is to come all resonated with past abuse. Some survivor participants said that they were even afraid of being abused by the health care practitioner:

[In the clinic waiting room, I felt] nervous, apprehensive, not exactly knowing what was going to happen ... as far as clothing was concerned or ... touch, just not knowing. (Woman survivor)

Discomfort with persons who are the same gender as their abuser(s). For some survivors, the gender of a person in a position of authority is a powerful “trigger” that can leave them feeling vulnerable and unsafe. This strong reaction prevents some survivors from seeking care from practitioners who are the same gender as their abuser:

[A male health care provider and assistant were] in the room with me, and I had my pants off, and this guy’s putting [ultrasound] gel on my leg. And I felt really uncomfortable ... even though ... probably nothing could have happened, but I just didn’t like the fact that I was in a room by myself with my pants off, with two men. That was really eerie. (Woman survivor)

My abuser was my mother. I don’t like to be touched by women, especially strange women. (Man survivor)

While it is understandable that health care practitioners have negative feelings in response to a patient’s negative transference, they must strive to contain these feelings and respond professionally.

Triggers. Examinations or treatments may “trigger” or precipitate flashbacks, a specific memory or overwhelming emotions such as fear, anxiety, terror, grief, or anger. A flashback is the experience of reliving something that happened in the past and usually involves intense emotion. Some survivors are particularly susceptible to flashbacks and some are overwhelmed by them:

And the goop that they put on me for the ultrasound gave me flashbacks, nightmares, insomnia; I just couldn’t deal with it. (Woman survivor)

Section 7.6 – Triggers and dissociation

Dissociation. Survivor participants also spoke about dissociating during interactions with health care providers. The Diagnostic and Statistical Manual of Mental Disorders, Fourth Edition (Text Revised) (DSM-IV-TR) explains dissociation as “a disruption in the usually integrated functions of consciousness, memory, identity, or perception of the environment” that may be sudden or gradual, transient or chronic. Some authors (e.g.,120,155) liken it to a state of divided consciousness in which aspects of the self that are normally integrated become fragmented. Dissociation is also understood to be a process that exists on a continuum,120 with one end being “common experiences such as daydreaming and lapses in attention, through déjà vu phenomena ... [and the other end of the continuum involving] a pathological failure to integrate thoughts, feelings, and actions.”111
Physical pain. For some survivors, the experience of acute and/or chronic physical pain may be associated with past abuse. This association can manifest itself in various ways (e.g., some individual have learned to ignore or dissociate from pain, while others are hypersensitive to it):

*I think sometimes when survivors are in pain, and coming for physical therapy, it hooks us back into...our childhood where we were in pain and...no one responded. And if you did indicate you were in pain...the pain was trivialized or you were threatened [so that you did not tell] anyone.* (Woman survivor)143p.256

**Ambivalence about the body.** Many survivors feel hate, shame, and guilt about their bodies. As children, many believed that something about them or their bodies invited or caused the abuse. This belief is reinforced if the survivor enjoyed some aspects of the abuse (e.g., special attention, physiological arousal).133 This shame and guilt may lead some survivors to feel ambivalent about and disconnected from their bodies:

*And [the amount of attention that I give to my body] ebbs and flows too, depending on where I’m at and how well I’m choosing to take care of my body. Which is a very difficult thing for me physically to do, because when you don’t live there, it’s just sort of a vehicle to get around.* (Woman survivor)143p.255

The conflict between the need to seek health care for a physical problem and the ambivalence or dislike of one’s body can affect treatment. For example, an individual may ignore symptoms that might contribute to an accurate diagnosis, explain an individual’s response to treatment, or
interfere with the ability to self-monitor effects of an intervention or medication.

**Conditioning to be passive.** Abuse can teach children to avoid speaking up or questioning authority figures. In adulthood, survivors may then have difficulty expressing their needs to a health care practitioner who is perceived as an authority figure.

[The health care practitioner did something and] I really freaked but ... I didn’t show her I was freaking, because our history is that you don’t let on if things are a problem for you. You just deal with it however you can ... by dissociating or what have you. (Woman survivor)

**Self-harm.** Self-harm (e.g., scratching, cutting, or burning the skin) is a way that some survivors attempt to cope with long-term feelings of distress. Health care practitioners may see evidence of self-harm in the form of injuries or scars on the arms, legs, or abdomen. Self-harm may take more subtle forms as well, such as ignoring health teachings or recommendations for treatment or symptom management (e.g., refusing to pace one’s activity in response to pain or fatigue, or failing to adhere to a diabetic treatment regime).

There are many reasons why survivors harm themselves. It may serve to distract them from emotional pain, focus the pain to one area of the body, or interrupt an episode of dissociation or numbness. Some survivors may harm themselves to regain a sense of control or ownership of their bodies. For others, it may be a punishment or an effort to atone for wrongs they believe they have committed. Dusty Miller argues that self-harm is one example of a range of self-destructive behaviours that can be thought of as an unconscious effort to reenact past trauma.

### 3.6 Questions about sexuality and sexual orientation

Survivors of child sexual abuse, like many other people in our society, may have questions about their sexuality or sexual orientation. Some male participants who had been abused by men said they had struggled with uncertainty about their own sexual orientation:

I just realized in sexual abuse, it seems very, very common that the issue of homosexuality when dealing with a male [survivor] of sexual abuse comes up. It’s an issue: “Am I a homosexual?” (Man survivor)

Some women survivors report similar struggles:

Female survivors of female-perpetrated abuse also experience this confusion around their sexual identity and orientation. (Woman survivor)

For participants who self-identified as gay, public assumptions about the “cause” of their sexual orientation and about their potential to be abusers were also problematic:

They assume that because it was your mother [who abused you] that’s why you’re gay. Because it was a woman doesn’t make much sense. Or that then because you’re gay, you were abused, you’re going to be a pedophile yourself. These attitudes come out from others that I’ve disclosed to. Lots of layers there; biases would be one of the big problems there with health practitioners. They’re going to make assumptions. (Man survivor)

Relatively few survivor participants raised the issue of sexual orientation in the context of their interactions with health care practitioners. However, a number of health care practitioner participants who commented on drafts of the Handbook pointed out that the phenomena of sexual identity and sexual orientation are often overlooked or ignored by health care practitioners. Certainly, it is important to recognize that
women and men who have been sexually abused in childhood may experience challenges around sexuality and intimacy in general. This is true of a proportion of survivors in heterosexual relationships as well as for some in same-sex relationships and for some survivors who identify as gay, lesbian, bisexual or transgendered.

Because of the general societal perception that being gay, lesbian, bisexual, or transgendered (GLBT) is “abnormal” or “wrong,” abuse survivors (and health care practitioners) may sometimes attribute their same-sex attraction to past sexual abuse. Shoshana Pollack, professor of social work at Wilfrid Laurier University, notes that “fostering this assumption in patients misses the important point that childhood sexual abuse involves traumatic sexualization and often leaves survivors confused about how to engage sexually in general, what their sexual preferences are (not only gender, but practices), what it means if they experience same sex attraction, what it means if they don’t experience it but as a child their abuser was the same sex etc.” (2007, personal communication).

No research has supported the idea that childhood sexual abuse is associated with the development of GLBT identity. No research studies have supported the claim that childhood sexual abuse is associated with the development of GLBT identity. In an online questionnaire study of lesbian and bisexual women between 18 and 23 years old, fewer than half of those who had experienced childhood sexual abuse thought that the childhood sexual abuse had affected their feelings about their sexuality or how they “came out.” Among those who did identify effects on their feelings about their sexuality and coming out process, some said that the abuse had not affected their feelings about their sexual orientation, which they believed was unconnected to the childhood sexual abuse experience.

Participants who were abused as children and who are (or have been) involved in same-sex relationships often have to deal with negative thoughts about themselves based on negative societal stereotypes. For example, some may think, “I’m bad because I was abused,” or “I am really bad because I was abused and it made me be attracted to the same sex.” These thoughts should be recognized as internalized heterosexist and homophobic social attitudes that need to be challenged and worked through. (Shoshana Pollack, 2007, personal communication)

The conflict between the need to seek health care for a physical problem and difficulty in caring for one’s body often affects treatment.
4 Principles of Sensitive Practice

4.1 Overarching consideration: Fostering feelings of safety for the survivor

*I now am beginning to understand that my physical wellness is really very connected to my emotional state, and if I'm not comfortable, if I'm feeling unsafe, then I'm not going to progress as quickly as [the health care practitioner] would want me to.* (Woman survivor)

The primary goal of Sensitive Practice is to facilitate feelings of safety for the client. The nine themes below were identified by virtually all participants as important to facilitating their sense of safety during interactions with health care practitioners. These themes are so critical to survivors’ feelings of safety that we term them the principles of Sensitive Practice. Through the course of our research, we have come to conceptualize safety as a protective umbrella, with the principles of Sensitive Practice being the spokes that hold the umbrella open. When the umbrella is open, an individual feels safe, and can participate in the examination or treatment at hand. While most of the principles are components of patient-centred care (see Stewart[63]), they take on even greater significance within the context of childhood sexual abuse and other interpersonal violence.

Child sexual abuse is a betrayal of trust and the antithesis of safety. Survivor participants frequently described to us how perpetrators, while abusing them, assured them that they were safe when just the opposite was true. For some adult survivors, the experience of being told that they are safe can trigger fear and anxiety. Thus it is clearly not enough for health care practitioners to simply assure their patients that they are safe. To facilitate survivors’ feelings of safety, practitioners need to make every effort to follow the principles of Sensitive Practice. To paraphrase one of the health care practitioner participants, the principles of Sensitive Practice articulate a standard of practice and provide a concrete and specific “how to” guide for doing this.

Since all health care practitioners – knowingly and unknowingly – work with individuals with histories of sexual, physical, and emotional abuse and other forms of violence, these principles represent a basic approach to care that should be extended to all clients. The principles of Sensitive Practice are analogous to the infection control guidelines (commonly termed “routine practice” or “universal procedures”) that have become part of everyday practice in all health care settings. Just as clinicians may not know an individual’s history of past infection, they may not know an individual’s abuse history. By adopting the principles of Sensitive Practice as the standard of care, health care practitioners make it less likely that they will inadvertently harm their patients or clients.

4.2 The nine principles of Sensitive Practice

**First Principle: Respect**

*[Feeling respected], to the person who has been abused, it certainly means a great deal.* (Man survivor)
Second Principle: Taking time

Time pressures – a reality in today’s health care system – constantly challenge clinicians to balance efficiency with good care. Sadly, this often leaves individuals feeling like one of many objects in a never-ending assembly line, and compounds survivors’ feelings of being depersonalized and devalued. For some, being rushed or treated like an object diminishes their sense of safety and undermines any care that follows.
Escalating patient-clinician ratios may lead many practitioners to become exclusively task-oriented, questioning whether they can afford the time to really listen to their patients. It is important to remember that feeling genuinely heard and therefore valued is healing in itself, and in some cases may be the most effective intervention a clinician has to offer:

Feeling genuinely heard and therefore valued is healing in itself, and in some cases may be the most effective intervention a clinician has to offer.

It’s the health care practitioners that ... stop and give you a moment, and that’s one of the biggest healing things right there, that moment. (Man survivor)\textsuperscript{159}

Sixth Principle – Respecting boundaries

Section 6.5: Time

Section 8.4 – Responding effectively to disclosure

Third Principle: Rapport

Showing some empathy, some caring, some concern ... make me feel that I’m a person as opposed to another client file going through. (Man survivor)

While rapport is essential to every therapeutic relationship, it is an absolute necessity to facilitate safety for survivors. Practitioners who are warm and compassionate facilitate good rapport and subsequent feelings of safety:

[For the health care practitioner I saw, this was] just a job like any other job. She could be answering phones. And I was just another name on a [referral] ... She wasn’t interested. She had no warmth ... I didn’t experience being safe with her because I didn’t think that this was somebody I could talk to at all, about anything! She just was not interested. (Woman survivor)\textsuperscript{143p.252}

Good rapport not only increases individuals’ sense of safety, but also facilitates clear communication and engenders cooperation. Survivor and clinician participants agreed that rapport is strengthened when clinicians are fully present and patient-centred.

The balance of professionalism and friendliness that contributes to positive rapport is partly a function of interpersonal style, but it can be developed with practice. Clinicians who are distant and cold in their professionalism are unlikely to facilitate a positive connection with clients. Conversely, an overly familiar style may be perceived as invasive and even disrespectful. Developing a tone that is professional and yet conveys genuine caring promotes a sense of safety and helps to establish and maintain appropriate boundaries.

Fourth Principle: Sharing Information

[He always gave] a reason why he was doing something, which was great ... It wasn’t just doing things and then leaving you in the dark. Or if he was asking questions, you don’t have to second guess, “Why did he ask that question?” Because my favourite sport is jumping to conclusions, right? ... If the person took ten seconds to tell me, “This is why I’m going to do it,” it will stop the mind from running. (Man survivor)

While knowing what to expect decreases anxiety for most people, it is particularly important for survivors. Survivor participants emphasized that they do not know what many health care practitioners do and therefore do not know what to expect. Being told what to expect on an ongoing basis helped to allay their fear and anxiety and often prevented them from being triggered by unanticipated events:

I think they should spend the five minutes at the beginning saying, “This is what [I] need to do to figure out what will best work for you,” so that we’re prepared, you know. The element of surprise is just really, really difficult to deal with ... and if there’s a preparation and there’s not that fear of the unknown, and
not the likelihood then that I will be triggered by something that is done, you know, into remembering something that is abusive for me.
(Woman survivor)\textsuperscript{143p.255}

Fifth Principle: Sharing control

A central aspect of sexual victimization is the loss of control over one’s body. It is understandable, then, that having a sense of personal control in interactions with health care providers who are more powerful is crucial to establishing and maintaining safety.

I’m learning that if I don’t have a sense of control ... I will walk away from [the situation].
(Woman survivor)\textsuperscript{143p.255}

Although both parties contribute to the dynamics of the helping relationship, the health care practitioner, by virtue of having greater social power and specialized training, has a greater responsibility in this area. Contracts for care, practitioner services contracts, and therapeutic contracts (either written or verbal) are all tools for articulating goals, clarifying roles and responsibilities, and defining the parameters of the helping relationship.

A frank, matter-of-fact discussion of these issues should be part of the treatment plan, as it serves to minimize miscommunication and misunderstanding and contributes to increased trust on the part of patient and health care practitioner alike. To proceed without such discussion assumes that clients and clinicians are all mind readers who, without deliberate effort, can clearly understand others’ words, motives, and intents.

Sharing control of what happens in the clinician-patient interaction enables individuals to be active participants in their own care, rather than passive recipients of treatment.

As the term \textit{sharing information} implies, it is a mutual process of information exchange in which both parties feel heard and understood.

Health care practitioners must also seek ongoing feedback about the patient’s reactions to the exam, treatment, or intervention throughout every encounter and prior to the next encounter. This invitation to articulate one’s reactions is particularly important for individuals who may indeed experience adverse reactions – such as flashbacks or nightmares – after an encounter. One man who had experienced oral abuse, for example, spoke about having nightmares for many successive nights after getting braces put on his teeth.

The surprises are the worst thing. (Man survivor)

In many cases, clinicians can begin the information sharing process before seeing the new patient by providing written information about what is involved in a patient appointment. Some clinicians share information by offering a running commentary on what they are doing as they are doing it. This does not require additional time, can be a tool for patient education, and is tremendously reassuring.

Appendix E – Sample Introduction to a Facility

As the term \textit{sharing information} implies, it is a mutual process of information exchange in which both parties feel heard and understood. A place to begin is to ask patients what information they want or need and to invite questions:

[The clinician] brings definite knowledge and expertise [into treatment] ... So together with what I know and what I can tell her, I would hope that she would be able to ... assess the situation and offer alternatives ... So instead of her being the expert and me being the patient, us being co-communicators about my body. That’s what I’d like to see.
(Woman survivor)\textsuperscript{144p.82}

\textbf{Appendix E – Sample Introduction to a Facility}

As the term \textit{sharing information} implies, it is a mutual process of information exchange in which both parties feel heard and understood.

\textbf{Sharing control of what happens in the clinician-patient interaction enables individuals to be active participants in their own care, rather than passive recipients of treatment.}
some other way that you feel more comfortable – help us, help us so that we can help you out. Let’s communicate here, let’s talk about things. I can’t read your mind ... I care enough about you to consult with you. To make you part of the healing process rather than a recipient.” You know? You need to be part of it. (Man survivor)

The process of ascertaining informed consent is a vital part of sharing control, as well as a legal responsibility. Informing, consulting, and offering choices are all part of seeking consent:

*It’s the approach for me. That immediate taking over, taking over for me without consulting me or giving me a choice ... For me that’s the first thing that raises my anxiety level ... for instance if you lay on a table, [the health care practitioner could say], “Are you okay to lay sideways or are you okay to lay on your back?”*, instead of telling me, “You lay on your back.” ... It goes back to education in a sense: “This is the procedure that we’ll be doing and this is what is expected of you.” ... So information and then choice. (Woman survivor)\(^{159}\)

By demonstrating respect for and sensitivity to personal boundaries, clinicians model healthy boundaries and reinforce patients’ worth and right to personal autonomy.

Section 6.6 – Informed consent

The health care practitioner must directly address all clients – even those who are minors, speaking through an interpreter, or cognitively impaired – and negotiate care with them.

**Sixth Principle: Respecting boundaries**

*As a survivor, I need to know that that person is not going to invade my space. Or do harm to me. Not necessarily physically, but emotionally.* (Woman survivor)\(^{164}\) p.95

Because respect for boundaries is crucial to a sense of safety for most survivors, it is a principle in its own right, separate from the first principle of Sensitive Practice, “respect.” The provision of health care often requires clinicians to work in close physical proximity to patients and to seek information of an intimate nature. Survivors said that health care practitioners’ questions and actions when initiated either without explanation or without permission left them feeling violated.

Violation of a client’s personal boundaries may occur unintentionally. For example, a practitioner, when rushed for time, may neglect to ask for consent before beginning a procedure. Although this action may meet the health care practitioner’s need for expedience, it does so at the expense of the client’s need for control and autonomy. Similarly, asking a very personal question before establishing rapport can be perceived as a psychological breach:

*My concerns when seeing a health care practitioner are related to the problems that I experienced as a child, and I’m still affected by them, and when somebody’s going to cross my – what I call my personal boundaries, the space that’s around me, that I call my own ... and if anyone else is coming into that space, I prefer that they tell me exactly what they’re doing there. When it comes to doctors, more so than anybody else, because they have a tendency to approach you ... with their hands out to go to work. I just can’t accept that because of the feelings and the stress and the emotions that are created in me are just too hard on me.* (Man survivor)

Learning about boundaries and boundary maintenance is a lifelong process. The blatant disregard of personal boundaries during abuse teaches children that their wants and needs are of little consequence. For many survivors, healing from abuse involves establishing or reestablishing personal boundaries and learning healthy and effective boundary maintenance strategies. By demonstrating respect for and sensitivity to personal boundaries, clinicians model healthy boundaries and reinforce patients’ worth and right to personal autonomy.

It is also possible that a clinician’s boundaries may be violated. For example, the patient who
they may need encouragement to become full, active participants in their own health care. Many of the survivors in our studies talked about the importance of even small encouragements from health care practitioners and of how they carry these encouragements into other life situations:

That assertiveness of [saying] no takes a long time to get ... it was somebody else giving me permission that allowed me to say no until I could learn to give myself permission [to do so]. (Woman survivor) 143p.254

I often need the “permission” later in the examination, when my trust has built, to be able to speak or ask about those things as well. (Man survivor)

As health care practitioners learn about the health effects of interpersonal violence and about working effectively with survivors, their best teachers will be survivors themselves.

Second Principle – Taking time

Boundary maintenance is a fiduciary responsibility clearly spelled out in professional codes of ethics, and violations carry serious sanctions. Addressing boundary problems in a direct, matter-of-fact way helps ensure patient safety and helps health care practitioners avoid potentially dangerous or compromising situations. While effective boundary maintenance may seem simple at first glance, it can be just the opposite and so requires the ongoing, lifelong attention of every health care provider. Practitioners who encounter specific difficulties are encouraged to consult with a respected peer or supervisor or seek advice from their professional body.

Seventh Principle: Fostering mutual learning

The principles of Sensitive Practice are intended to increase clients’ sense of interpersonal safety. Because many of them have not experienced that sense of safety as children, abuse survivors may be learning about it only in adulthood. Thus, compassion and interest to learn about the health effects of interpersonal violence and about their particular needs. In the context of a caring relationship, most survivors are even willing to tolerate missteps and the inevitable discomfort that comes with addressing a difficult topic.

Eighth Principle: Understanding non-linear healing

Survivor participants reminded us repeatedly that healing/recovery from childhood sexual abuse is not a linear process. As a result, the degree to which a survivor is able to tolerate or participate in treatment may vary from one health care encounter to the next. This variability may occur over the short term (day to day) or over longer periods of time. In recognition of this reality, health care practitioners must check in with their clients throughout each encounter and adjust
their behaviour accordingly. The practitioner who responds with understanding and compassion in these circumstances contributes to the survivor’s feeling of safety and to a stronger therapeutic alliance:

\[\text{Part of my body at different times might be untouchable. It’s going to change, depending on what I’m dealing with. So, you’re not going to be able to make a list and count on that every time … it’s going to be a check-in every session.} \text{(Woman survivor)}\]

**Ninth Principle: Demonstrating awareness and knowledge of interpersonal violence**

[The health care practitioner] had a book and a pamphlet on a table nearby where I was sitting that talked about sexual abuse, and so immediately that said to me, number one, she is open to this and therefore if it comes up I know that I’m in good hands because [otherwise] this stuff would not be sitting here. \text{(Woman survivor)}

Many survivors look for indicators of a clinician’s awareness of issues of interpersonal violence. Evidence of this awareness can take a variety of forms. Posters and pamphlets from local organizations that serve those who have experienced violence may help a survivor overcome hesitancy in raising the issue with a health care practitioner. In addition to an indication that their health care practitioners have an understanding of interpersonal violence, male survivors may also be looking for an indication that they are aware that men may be survivors:

\[\text{A poster in all the examining rooms. You know – victims of child abuse are welcome. That’s easy. Male victims of child abuse validated here. We care about the victimization of children, help prevent victimization of male children. Those are messages that you can put on posters. Let’s protect little boys and girls – see, inclusive. Boys and girls who have been victimized as children are welcome. Boys and girls … [and] have the picture – boy and girl.} \text{(Man survivor)}\]

Incorporating the principles of Sensitive Practice into daily practice also indicates a health care practitioner’s awareness of issues related to interpersonal violence.

### 4.3 Using the principles to avoid retraumatization

The nature and quality of the relationship between a clinician and a survivor has implications for the safety and effectiveness of health care. A good helping relationship not only contributes to an open exchange of information, but also creates the “human-to-human” environment that is essential for the establishment of trust. Effective helping relationships are not ethereal, mystical connections that “just happen,” nor are they a naturally occurring byproduct of a charismatic personality. Effective helping relationships are intentional and skill-based interactions that exist to serve the needs of the patient. Effective helpers are genuine, empathic, and warm. They are also open-minded, knowledgeable, attentive to verbal and nonverbal communication, self-aware, and reflective.

Consciously applying the principles of Sensitive Practice can not only enhance the therapeutic relationship with the survivor but also assist the practitioner to avoid retraumatizing the patient. Many survivors spoke about how interactions with health care practitioners had left them feeling violated and retraumatized:

\[\text{It’s critical that they understand that we can be retraumatized as a result of how we are treated by them … Not that they’re meaning to go there, but by not treating us respectfully – giving us what we need to feel safe, and being allowed to be seen as co-partnering and not as having no power at all – [they are making it] possible for us to be retraumatized.}\]
And I would like them to get the information. (Woman survivor)\textsuperscript{159}

Without attention to these principles, survivors’ umbrellas of safety can collapse, interfering with their ability to benefit from or perhaps even tolerate health care interventions. Survivors told us repeatedly that this applies in all health care settings, including offices (physicians, dentists, chiropractors, massage therapists, naturopathic doctors, physical and occupational therapists, etc.), acute care hospitals, community-based care, long-term care, and rehabilitation settings.

4.4 Questions for reflection

- How willing am I to share control with my clients?

- What are my own personal boundaries? How do I know if they are being violated? Could any of my actions be seen as boundary violations by clients?

- How do I balance the demands of my whole practice with the need to take adequate time with each client?

- What might get in the way of communicating my respect for my patients?

- What is my own personal style of interacting with clients? Does it seem to foster rapport? Do I put effort into maintaining rapport with each patient over time?

- How do I ensure that patients have received what they feel is adequate information about examinations, treatment options, and treatment processes?
5 Guidelines for Sensitive Practice: Context of Encounters

5.1 Administrative staff and assistants

The quality of interactions with administrative staff and assistants who work in health care environments can affect survivors' feeling of safety. Participants overwhelmingly agreed that, in an office environment, their interactions with administrative staff and assistants set the tone for the practitioner-patient relationship. For these reasons, staff need to have some understanding of the dynamics and long-term effects of interpersonal violence and require coaching in applying the principles of Sensitive Practice in ways that will work in their specific environments.

In both hospitals and community-based settings, routines and procedures have evolved to be cost-efficient and to maximize the clinician's time. They may, however, be experienced as more clinician-centred and less patient-centred. In many offices, for example, it is common practice for the receptionist to ask about the nature of the problem in order to book the appropriate type and time of visit. Many survivors said they experienced this as an invasion of privacy, especially when they are seeking assistance with psychosocial or mental health problems. The clinician participants in our working groups suggested that a preferable approach would be for receptionists to ask whether the appointment was for a discussion or an exam. Office personnel who usher individuals to examination areas and carry out preliminary procedures could also demonstrate respect for privacy by using this kind of question. Health care practitioner participants also reminded us that assistants and technicians (such as physical therapy assistants or x-ray technicians) who work directly with patient evaluation and treatment should use Sensitive Practice in the same ways that the clinician does.

One survivor emphasized the need for receptionists to learn about Sensitive Practice when she described her attempt to make an appointment with her family physician, who had previously agreed to see her if she was feeling suicidal:

*What do I have to do, stand up on a chair and say, “Yes, I look fine but at this moment I am thinking of a thousand and one ways to kill myself”? [When the receptionist refuses to give me an appointment] the shame and guilt kick in and I blame myself and I do go home and I OD or I slash my wrists.* (Woman survivor)

5.2 Waiting and waiting areas

Survivor participants spoke at length about the extreme anxiety that they experience while waiting for health care appointments because it takes them back to past abuse experiences. Because of their naivety, children never anticipate the first episode of abuse; it catches them unaware and defenceless. The sexual acts seem strange and may be painful; the secrecy is confusing; and the coercion or threat of harm is frightening. Children have no prior reference from which to understand why someone, especially someone they love and trust, would do these things to them. After the abuse has happened once, many children are haunted by the fear that it could happen again. They become hypervigilant and watchful, and wait in dread for the abuse to reoccur.

Although waiting for appointments is a fact of life, the experience may be particularly trying for survivors who have never completely shed the apprehension associated with waiting. Therefore, participants urged practitioners to:

- Create waiting areas that are warm and welcoming;
- Provide and clearly identify washrooms;
- Create waiting areas that are warm and welcoming;
• Provide printed materials related to interpersonal violence;

• Provide a realistic estimate of the length of wait time.

5.3 Privacy

Privacy is another important environmental aspect of survivors’ feelings of safety. The balance between safety and privacy is not the same for all survivors: some will be most comfortable in a private room; others may choose this option as long as they can be accompanied by a support person; and still others feel safer in public spaces. Many survivor participants ask that health care providers approaching a waiting client knock or announce themselves and await permission before entering.

Specifically, clinicians should consider the privacy (or lack thereof) that their practice environment affords, by reflecting upon the following questions:

• What can be heard and seen in the reception area?

• Are patients required to respond to personal questions in a public reception area where others may overhear the exchange of information?

• What can be heard and seen from the hallway?

• What can be heard between examination rooms or cubicles?

Practitioners are further urged to have at least one soundproof examination or interview area available for use. While privacy is even more difficult (and sometimes impossible) to achieve in hospital settings, clinicians are urged to be creative and to consider the possibility of using areas that are not soundproof when others are not present, such as during mealtimes and outside of peak hours.

5.4 Other issues related to physical environment

Having had so little control over what happened to them in childhood, many survivors seek ways to control the current physical environment in order to feel safe. They offer the following comments and suggestions:

• Designate separate washrooms for men and women;

• Take the time to familiarize the client with the physical environment (e.g., waiting area, washrooms, patient care areas, equipment, and emergency exits);

• Whenever possible, offer clients a choice of where they may sit in examination, treatment and waiting rooms (e.g., some survivors prefer to sit near or be able to see the door);

• Because some survivors are strongly affected by lighting and views of floors and ceilings, ask clients about their comfort level with the lighting. This is particularly important if treatment requires the patient to assume a position facing the floor or ceiling;

• For practitioners who use music, candles, or scent during treatment, check with the patient regarding their preference to avoid triggering negative responses.

5.5 Patient preparation

The importance of sharing information about health care procedures prior to beginning any exam, treatment, or hospital admission cannot be overstated. Since advance preparation can help significantly to reduce an individual’s anxiety even before the clinician and client meet, practitioners should consider the following possibilities:

• Send printed information to clients before their first appointment or give it to them...
5.6 Encouraging the presence of a support person or “chaperone”

[The presence of the assistant] would make me feel more comfortable if the door had to be closed ... it wouldn’t be that one-on-one. (Man survivor)

A third party observer (either a patient-nominated support person or clinician-nominated “chaperone”) is commonly used for some examinations and procedures. Survivors explained that having a support person with them often helps to decrease their fears. The support person can also serve as another set of ears to hear any information offered by the clinician:

If you’re being given a lot of information and you can’t necessarily hold it or get it all straight or if you’re anxious ... and there’s a lot of new information coming at you, it’s nice to have somebody in the room that can help you remember what’s being said. (Woman survivor)

The presence of a support person requires balancing competing demands for confidentiality, support, and protection of both patients and practitioners. To facilitate both the patients’ abilities to take advantage of the option of having a support person present and the integrity of their practices, health care providers are encouraged to:

- Inform patients verbally as well as via brochures and signs in the waiting areas about the option of having a third party observer with them;
- Remain aware that the presence of a support person may not always be in the client’s best interests (e.g., a violent partner who seeks to control the client’s interactions with others);
- Speak privately with the client at the beginning of the appointment to ensure that the individual actually wants to have

while they wait for their first appointment. Also consider displaying it in waiting rooms or treatment areas. These materials should be written in clear, plain language that avoids jargon or medical terminology. As well as providing information about the organization and service, these materials can also cue survivors to think about what they can do to facilitate their own safety (e.g., bringing a support person or a small familiar object that symbolizes safety and security with them to appointments).

Appendix E – Sample Introduction to a Facility

- Because not all clients are able to read written material or understand English, alternate strategies can be used to inform them about what they can expect in the health care encounter (e.g., consider using drawings, photographs, or videos that answer frequent questions and explain what will happen from the beginning to the end of the interaction).

- Helping any client prepare for hospitalization or outpatient procedures begins by assessing what they know and identifying any knowledge gaps. Responding to those gaps may involve brainstorming and negotiation as well as information sharing. For example, when working with abuse survivors it is important to discuss: (a) ways that the survivor can get through the experience in the least traumatic way; (b) ways to avoid identifiable triggers; and (c) plans to ensure sufficient ongoing support.

Collaborating with clients to develop a written plan of care ensures that everyone who works with them is aware of their particular needs.
A critical cultural perspective, and understanding culture as relational, shifts the gaze away from cultural Others onto the self, and requires examination of how each individual is enmeshed within historical, social, economic, and political relationships and processes. This then leads to questions such as: How am I reinforcing certain norms (for example, Eurocentric norms perhaps) within the culture of health care? How am I seeing certain behaviours, beliefs, and practices as “normal” and others as “cultural”? How am I serving certain economic and political interests through my daily practices?

These ideas are also reflected in Irihapeti Ramsden’s work on cultural safety. Ramsden, a Maori nurse leader in New Zealand, developed the concept of cultural safety to bring attention to the negative impact of colonization on the health of Maori people and the ways in which colonization privileged Eurocentric health/illness beliefs and many current practices perpetuated inequalities. A full discussion of this topic is beyond the scope of this handbook; however, health care practitioners and health care organizations are encouraged to examine current practice with diverse groups on an ongoing basis.

While interpersonal violence is present in all cultural and ethnic groups, we pay special attention to Aboriginal Peoples in this Handbook because they represent a significant and growing portion of the Canadian population, they continue to experience the long-term effects of widespread abuse in residential schools, and, in our view, are likely to benefit from the universal application of the principles of Sensitive Practice. It is our hope that even a basic understanding of the abuses (many of them systemic) that large numbers of Aboriginal people experience will help health care practitioners be more sensitive and therefore more effective in health care interactions with Aboriginal peoples.

5.7 Working with survivors from diverse cultural groups

Because Canadian society is composed of individuals from many racial, ethnic, and cultural groups, it is imperative that health care be culturally sensitive. Although much has been written about cultural awareness, cultural sensitivity, and cultural competence in health care, ideas about how to address the topic of culture are continuing to evolve. Early work in the area (for example,37,38,19,20,21) offered models of cultural competence as a framework for delivering responsive health care services to individuals from culturally and ethnically diverse backgrounds. More recently, proponents of the critical cultural perspective (e.g.,34,74,75) have encouraged practitioners to broaden their thinking about culture beyond that described in these early models and to recognize culture as a complex, dynamic, and relational process that is shaped by historical, social, economic, and political forces. As University of British Columbia nursing professors Annette Browne and Colleen Varcoe explain,
5.8 Collaborative service delivery

Survivors spoke about instances in which they did not feel comfortable or safe working with health care providers to whom they had been referred. While options may be limited, all patients have the right to referral to another clinician or facility. Clinicians may also find themselves in a position in which they cannot meet survivors’ expectations or needs for care. Broaching such discussions may not be easy, but practitioners are encouraged to respond to requests for referral and to be supportive in discussing situations in which they believe they cannot meet patients’ needs.

The transfer of care from one practitioner to another without prior notice can evoke feelings of abandonment and erode trust. Whenever possible, individuals should be offered a choice of alternate caregivers. Announcing planned absences well in advance provides clients with the option of making alternative arrangements. In the best possible scenario, clinicians are able to introduce their clients to the practitioner who is taking over. Discussion about what information regarding past abuse the individual consents to be given to the temporary caregiver is essential. For permanent transfer of care, the “outgoing practitioner” should ensure that the new colleague is knowledgeable about interpersonal violence and the sensitive care of survivors.

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Survivors urged health care providers to consider making (with permission) informal links with other practitioners with whom they were working to address health problems more fully.

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As a survivor of abuse, [I feel that today’s health care system] is reobjectifying ... to the point where I scarcely exist ... as a whole being because society [has] modeled a dissociative process that took my emotions to a psychiatrist, and my body to a GP, and my teeth to a dentist. They didn’t show me any model that would pull me back out of dissociation. (Man survivor)

Survivors agreed that they did not expect any one person – including a health care practitioner – to fix all of their problems. Some suggested that access to a range of practitioners from different health care disciplines would be an asset to their healing and those who had experience with primary health care teams were very positive about that experience.

Many participants were aware of the connections between mind, body, and spiritual wellbeing and wished that health services were more holistic in their approach:

I think that the connection between mental health and physical health can’t be separated ... The [practitioner] would be able to help a client deal with their health issues significantly more if they understood what the underlying emotional stuff was as opposed to never, ever asking the question and possibly figuring this out ... I think that the role of the [practitioner] in health should [include] a larger component of emotional health ... I think that I ... could have got to the place of dealing with the emotional place and impacts of sexual abuse an awful lot sooner if there had been some help to sort of draw that out. (Man survivor)

Guidelines for Sensitive Practice: Context of Encounters
example, a conversation between a counsellor and a practitioner about a treatment that a survivor experiences as triggering intense negative emotion might lead the counsellor to work with the survivor on grounding techniques and to offer the practitioner additional suggestions to minimize these reactions:

*I was quite amazed and thrilled that I could go in to see my [psycho]therapist and ... during the week she and the medical doctor and the psychiatrist had talked about my case and, you know, they were all concerned on a certain level about a certain thing.* (Man survivor)

5.9 Practitioners’ self-care

Taking care of oneself – eating well, getting enough rest, engaging in regular physical activity, taking time to relax, and so on – can be a challenge for most people. For survivors who learned as children that their needs are not important, self-care may be even more difficult:

*This is the first time in my life for the past three years that I've given a damn about my physical well-being. I never gave a damn before. That's due to living with very poor self-esteem.* (Man survivor)

An important aspect of health teaching is the modelling of self-awareness and self-care. Patients who have difficulty in these areas may learn from seeing their health care practitioners modelling self-care and appropriate boundary setting.

Section 4.2 – Sixth Principle: Respecting boundaries

The basic tenet of self-care for practitioners is the need to extend to themselves the understanding and compassion that they demonstrate to their patients. Every clinician needs to develop and use a repertoire of strategies that promote and maintain health, particularly during stressful or emotionally intense encounters with patients. It is also crucial to remember that the capacity to work through difficult situations is never constant, even for experienced practitioners.

Practitioners may need to seek the support of a colleague or counsellor to talk about their own reactions to disclosures of childhood sexual abuse or other difficult situations with patients. Obtaining this support can and must be done without breaching confidentiality. Seeking support is not a sign of weakness; rather, it is indicative of taking professional responsibilities seriously. Ignoring one’s distress or discomfort increases the risk for Secondary Traumatic Stress Disorder (STSD), also known as Vicarious Traumatization (VT), or Compassion Fatigue (CF). Charles Figley, director of the Traumatology Institute at Florida State University, described the symptoms of STSD as being similar to those of Post-Traumatic Stress Disorder (PTSD) "except that exposure to a traumatizing event experienced by one person becomes as traumatizing even for the second person." Individuals with PTSD or STSD may experience depression, anxiety, lethargy, overinvolvement with abused patients, and undue fear of personal and familial abuse. If these symptoms go unrecognized and untreated, practitioners may react by avoiding abused clients or inadvertently conveying to them that they have done something wrong.

For health care practitioners who are also survivors. It is also important to keep in mind that childhood sexual abuse survivors and health care providers are not categorically discrete groups. A proportion of clinicians are themselves survivors of childhood sexual abuse. Practitioners who have personal histories of childhood sexual abuse may be especially empathic towards other survivors, particularly if they have worked through and resolved their own wounds. However, practitioners who have unresolved abuse issues may face great challenges when working with other survivors. They may be at risk for being triggered, developing boundary problems, and counter-transferring harmful responses to patients. It is recommended that individuals work through and come to terms with their own history of childhood sexual abuse to
avoid confusing their own difficulties with those of their patients.

5.10 Community resources for survivors and health care practitioners

Survivors clearly do not expect health care practitioners to be all things to all people. At the same time, practitioners can play a vital role in helping their patients locate and access appropriate services and resources. Organizations such as sexual assault centres, women’s centres, community mental health agencies, and residential addiction treatment facilities may provide information for survivors and practitioners. Organizations serving male survivors have emerged in some communities in recognition that many organizations established earlier were serving only women. Many sexual assault centres can offer specialized training or support for clinicians in their work with childhood sexual abuse survivors. Practitioners in the community who have expertise in working with survivors may be available, and they may be willing to consult or mentor other health care providers. Professional associations and regulatory/licensing bodies may be able to suggest other available resources.

Gathering information on the following questions will help practitioners determine whether an organization is appropriate for counselling referrals:

- The agency’s mandate and the nature of services offered (e.g., crisis intervention, individual counselling, group therapy, support groups).
- The agency’s policy on fees for service (e.g., what the fees are, whether the agency offers a sliding scale, whether it accepts payment from second parties such as employee health plans).
- How soon a prospective client can expect to receive service and whether there is a wait list.

Prominently displaying posters and brochures for programs and agencies that serve survivors of interpersonal violence offers patients the message that the practitioner is aware of the prevalence and potential long-term problems associated with sexual, physical, and emotional abuse. Materials should provide information on:

- Sexual assault centres, women’s centres, community mental health agencies, and residential addiction treatment facilities;
- Telephone help lines and suicide hotlines;
- Battered women’s shelters;
- Mobile crisis units.

Prominently displaying posters and brochures for programs and agencies that serve abuse survivors offers patients the message that the practitioner is aware of the prevalence and potential long-term problems associated with sexual, physical, and emotional abuse. Materials should provide information on:

Recommended Reading and Resources
6 Guidelines for Sensitive Practice: Encounters with Patients

6.1 Introductions and negotiating roles

In all health care settings, steps must be taken to ensure that the first moments of an encounter set a tone consistent with Sensitive Practice. By introducing oneself, explaining the nature of the appointment, and asking patients how they wish to be addressed, practitioners convey respect for their clients and begin to build a positive relationship with them.

Further, before beginning any intervention, health care providers must ask clients about their expectations for care. Doing so establishes a relationship that involves two-way sharing of information and control. It also creates an opportunity for the clinician to gain quick insight about potential apprehensions, which can help to avoid triggering negative reactions. In long-term health care relationships, the periodic revisiting of roles and responsibilities allows for renegotiation and communicates genuine compassion and concern.

Steps must be taken to ensure that the first moments of an encounter set a tone consistent with Sensitive Practice. 

While the standards of care for certain examinations require removal of clothing, survivor and clinician participants alike urged all health care practitioners to consider the following:

- Discuss clothing requirements with patients and collaborate with them to find an agreeable solution (e.g., allowing clients to wear their undergarments throughout the exam or inviting clients to wear their own abbreviated clothing such as bathing suits or shorts).
- Leave the room while the patient is changing.
- Provide a variety of sizes of gowns for all body sizes and instruct the patient about whether the opening is to be at the front or back.
- Avoid paper gowns whenever possible (they were widely described by survivors as so flimsy that they escalate feelings of vulnerability).
- Do not assume that all men are comfortable baring their chests.
- Meet patients when they are fully clothed (e.g., to make contact, ascertain the reason for their visit, or perform a health history).

If I had to take off clothing ... for a male [clinician] it’s ... hard because there’s the trust issue there and for me there was a lot of guilt and shame ... I struggle with body image and sometimes ... I feel powerless then. (Man survivor)\textsuperscript{159}

6.2 Clothing

Few issues highlight survivors’ difficulties in health care settings as much as the need for removal of clothing. While practitioners often take for granted the need to disrobe and to don an examination gown, undressing for someone in a position of authority transports many survivors back to their abuse and leaves them feeling powerless, vulnerable, and filled with shame:

Undressing for someone in a position of authority transports many survivors back to their abuse and leaves them feeling powerless, vulnerable, and filled with shame.

Section 4.2 – Fifth Principle: Sharing control
If clients need to disrobe for an examination or procedure, explain what will happen next, what level of undress is required, and why. Before proceeding, ask whether the client agrees to proceed with what has been explained and ensure that the client’s questions have been fully answered.

- Expose only the body area necessary for the specific intervention at any one time.

- Cover clients’ bodies as soon as exams are completed to minimize the length of time that they are exposed.

- Meet with clients again once the examination or procedure is finished and they have re-dressed to offer health teaching, provide an opportunity for questions, and say goodbye. While this may take a few extra minutes, it brings closure to the interaction and allows the client to leave on equal footing.

### 6.3 Task-specific inquiry

A task-specific inquiry involves asking patients about their preferences for or potential difficulties with a specific examination, procedure, or treatment. It provides an opportunity for patients to offer health providers information that is directly pertinent to the present situation without any reference to past interpersonal violence. Task-specific inquiry should be used during an initial meeting with a patient, before any new examination or procedure, and any time body language suggests that the patient may be uncomfortable or experiencing difficulty. Regardless of other factors, it should also be done intermittently during interactions as an ongoing invitation to offer feedback or to identify problems.

Task-specific inquiries involve a combination of closed- and open-ended questions that offer patients an opportunity to share anything that they consider relevant. A closed-ended inquiry might be, “Have you ever had difficulty with examinations/procedures like this one?” If the individual answers in the affirmative, then an open-ended question – such as “What can I do to make this easier for you?” – can help to minimize the patient’s discomfort. Before the examination begins, extend a broad invitation to share relevant information (e.g., “Is there anything else I should know before we begin?”).

Health care providers should recognize that, while asking for this information may seem safer than talking about past abuse, survivors may still experience it as difficult. If an individual’s nonverbal cues indicate tension or anxiety, the practitioner may need to ask very specific questions such as, “Do you have any discomfort having your blood pressure taken?” or “Do you have difficulty when someone touches your knees?” Survivor participants urged clinicians to:

> Pick up on obvious things: “You seem very anxious, is there something that you are uncomfortable with or is there something I should know?” (Man survivor)

There are many reasons that people experience specific discomforts and sensitivities during health care encounters – some of these relate to past abuse, but others do not. Therefore, while task-specific inquiry should be used for all patients, clinicians should be careful not to assume that a patient who offers a task-specific disclosure is an abuse survivor.

Task-specific inquiry should be used during an initial meeting with a patient, before any new examination or procedure, and any time body language suggests that the patient may be uncomfortable or experiencing difficulty. Regardless of other factors, it should also be done intermittently during interactions as an ongoing invitation to offer feedback or to identify problems.

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**Task-specific inquiry should be used during an initial meeting with a patient, before any new examination or procedure, and any time body language suggests that the patient may be uncomfortable or experiencing difficulty.**

**Asking if [the individual] has any issues or any concerns or are they uncomfortable, either physically or emotionally, is a really good way to start.** (Woman survivor)

One woman suggested that clinicians:

> Start out with, “What are your experiences with a dentist [doctor/massage therapist etc.]? How often do you go? What are your fears?” (Woman survivor)
Inquiries about sensitivities, discomforts, and difficulties can also be included on questionnaires that are part of an initial assessment. Some survivor participants told us that they are more comfortable with this approach. The practitioner may invite an individual who indicates having difficulty with a number of components of an examination to outline the issues in writing. Regardless of the mode used to elicit this information, it is vital to the sensitive health care of all clients.

Practitioners also need to be aware that, although they should make task-specific inquiries prior to any examination, some individuals may not be able to talk about their difficulties until they develop a rapport with their health care provider. Further, the ability or willingness to talk about task-specific difficulties may be a function of the survivor’s stage of healing; certain components of an exam may be well tolerated at some times and problematic at others.

Clinicians should not assume that an individual has disclosed all task-specific difficulties during previous interactions. Survivors who have been conditioned to be passive or to defer to authority may need ongoing permission and encouragement to talk about difficulties on a regular basis. Body language – such as trembling, flinching, tensing muscles, changing breathing patterns, flushing, crying, or dissociating (i.e., appearing spaced-out, distant, or blank) – should be explored as cues signalling that an individual may be experiencing difficulty:

We send out signals ... to people that we have been abused ... I was sending signals out, and I don’t think the people were listening really and picking up on them ... [I would] cringe and move and I often said “What are you doing?” (Woman survivor)

When these cues are evident, practitioners should explain that the exam or procedure will be easier to complete if the patient can relax, and then ask for assistance in discovering another approach to that component of the exam (e.g., “Would it help if I gave you a mirror to help you to see what I am doing?”). Addressing an individual’s apparent discomfort in these ways is vital to establishing and maintaining rapport; ignoring these things can undermine feelings of safety and trust.

Body language should be explored as cues signalling that an individual may be experiencing difficulty.

Task-specific inquiry should not be reserved exclusively for examinations involving touch.
Documenting task-specific difficulties or preferences can be done in a way that identifies the sensitivity or difficulty without any reference to past abuse. In deciding what to include in the patient’s record, practitioners should ask themselves what other clinicians need to know in order to provide the best care. When practitioners learn about task-specific sensitivities for the first time only after a patient experiences an adverse reaction to part of an exam (or treatment), they should record the unexpected response as soon as possible after the incident, including both objective (who, what, when, where, how, and how much) and subjective (what the patient and others report about the event) information.

### TABLE 2
A brief summary of task-specific inquiry

Task-specific inquiry involves asking patients about their preferences for or potential difficulties with a specific examination, procedure, or treatment. It provides an opportunity for patients to offer health providers information that is directly pertinent to the present situation without any reference to past interpersonal violence.

- Use combination of closed- and open-ended questions to offer patients an opportunity to share anything they consider relevant.
- An initial closed-ended question such as:
  - “Have you ever had difficulty with examinations/procedures like this one?”

If the individual answers in the affirmative, follow-up using an open-ended question such as:

- “What can I do to make it easier for you?”

- Before beginning an exam, offer one additional opportunity to disclose something the patient thinks might be relevant:
  - “Is there anything else I should know before we begin?”

- When a clinician notes discomfort, return to task-specific inquiry. For example,
  - “Every time I go to stand behind you to take a look at your back you seem to tense up. Do you have difficulty with having someone standing behind you or touching your back?”

If the patient responds in the affirmative:

- “What can I do to make this part of the exam easier for you?”

- For survivors who verbally deny discomfort but whose body language suggests the opposite, task-specific inquiry is likely to facilitate feelings of safety for the survivor; explain that carrying out the exam with the patient’s body more relaxed is ideal, and ask the patient’s assistance to discover another approach to that component of the exam. For example,
  - “Would it help if I gave you a mirror to help you see what I am doing?”

- Documentation of task-specific disclosures can be done in a way that identifies the sensitivity or difficulty without any reference to past abuse. Focus on communicating the information that will assist other healthcare professionals to avoid the difficulties or to problem solve with the patient to minimize the difficulties.
6.4 General suggestions for examinations

In keeping with the principle of information sharing, it is important that health care providers not assume that their patients know what they are doing or understand why they are doing it. Thus, it is important that they provide a running commentary of an examination or procedure explaining what is being done and why. Further, repeated invitations for questions are crucial:

*I found quite often when you go to a health care practitioner, they automatically assume that you have some kind of knowledge of their job outline ... And why should I know? I didn’t go to school for that, so it’s really frustrating. And they expect you to know something about it.*

(Woman survivor)143

While interactions with clients may be routine by clinicians, for many survivors, health care environments are strange and frightening places. Simply being in such environments can challenge an individual’s ability to ask questions or to verbalize their needs. This fact cannot be emphasized strongly enough:

*I feel very (pause) almost frightened. To some degree the fight or flight syndrome kicks in where I’m ready to hit the floor and head for the door. I know it’s because of problems that I’ve had as a kid. But at the same time those feelings come over me and then I lose all train of thought as to what I’m there for, what I want to ask him, I forget what day it is.*

(Man survivor)

While employing the following strategies for sensitively conducting an examination or treatment may require extra time, they are important for establishing rapport, trust, and safety – and they may actually save time in the long run:

- Complete the initial health history before asking the client to remove any clothing required for the physical examination.
- Invite patients to make a list of questions and concerns for each future appointment in order to reduce their sense of anxiety.
- Encourage questions throughout the encounter.
- Allow enough time to help individuals understand fully what is being done.
- Seek a balance between offering descriptors of symptoms (“Would you describe the pain as sharp or dull, throbbing or aching?”) and encouraging survivors to find their own words. This strategy can be particularly important since many survivors have learned to ignore their bodies and may require extra time to describe their symptoms.
- Move on from topics that are making individuals seem uncomfortable or questions that they are having difficulty answering, and return to them later.
- Use a written Informed Consent Form that uses readily understandable language and avoids abbreviations, jargon, and technical terms.
- Inform patients that consent can be withdrawn at any time, without penalty to them.
- Pay close attention to language to ease the patients’ anxiety (e.g., ask patients to change rather than to get undressed, use the term examination table rather than bed, and use the terms underwear or undergarments instead of panties).

Section 6.6 – Informed consent

Appendix F – Using Plain Language in Consent Forms (including a sample)

Recommended Readings and Resources – Plain Language: Websites
6.5 Time

- Most of the survivors we spoke with recognize the time pressures under which health care practitioners work and do not expect exceptional treatment. The following three suggestions are offered by survivor and health care practitioner participants as ways to use the available time more effectively:

- Inform clients at the outset of an appointment/interaction how much time you have to spend with them and negotiate how best to use it. In a clinic setting, a health care practitioner might say, “We have 15 minutes for this appointment, what do we need to focus on?” or “We have 15 minutes and I plan to ... Is there anything else that you need or want?” A health care practitioner in a hospital emergency department might say, “My name is ... I am a Registered Nurse and will be looking after you. We are very busy here today, so you may have to wait an hour or more to see a doctor.” The nurse might go on to say, “I need to get some information from you and then I’ll be in and out every 15 minutes or so to check on you. Please use the call button if you need me.”

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- Do not approach patients from behind and, because some individuals startle easily, avoid quick, unexpected movements.

- Notify patients before shifting focus from one area of the body to another.

- Explain the rationale for examining areas of the body other than the site of the symptom.

- Encourage individuals to ask that the examination or treatment be paused, slowed down, or stopped whenever it is necessary to lessen their discomfort or anxiety.

- Inform patients when procedures/examinations are likely to be uncomfortable and collaborate with them to minimize the discomfort by soliciting and responding to feedback throughout the procedure (e.g., ask, “How are you doing? Can we continue?” and, if they say “No,” take a break until they can continue).

- Avoid glib or false assurances which sound dismissive or indicate lack of understanding of their concerns (e.g., instead of saying “Trust me” or “Don’t worry, you’ll be fine,” say “I know this is difficult for you. How can I help you to feel more comfortable?”).

Even though employing these strategies may require some extra time, they are important for establishing rapport, trust, and safety - and they may actually save time in the long run.
to take telephone calls without any explanation, the message to patients is that the practitioner is not focused on them.

Throughout this project, we have heard repeatedly that short-term interactions pose the greatest difficulty for integration of Sensitive Practice because of the reality of time pressures. Survivor and health care practitioner participants alike urged that while it may take some commitment and ingenuity to incorporate the principles of Sensitive Practice in short-term interactions, it is important to make the effort to do so.

6.6 Informed consent

Obtaining informed consent for examination and treatment is an important part of practitioners’ responsibility to their clients and is regulated by professional/licensing bodies as well as legislation. Clinicians are urged to ensure that they are thoroughly familiar with all appropriate sources of information about informed consent. Informed consent involves: (a) explaining the health problem; and (b) making recommendations for addressing the problem (which must include: a discussion about the nature, benefits, material risks, and side-effects of treatment; alternative courses of action; and likely consequences of not having the treatment). Written consent forms are part of the process of obtaining informed consent in many health care settings and health care providers are urged to draft written consent forms in plain language.

This section highlights aspects of informed consent that are particularly pertinent to survivors. Because survivors have had early experiences of boundary violation, it is essential that practitioners be particularly attentive to obtaining consent which goes beyond the standardized forms and which is an ongoing aspect of their work with patients:

*When I was a child ... you might say yes [consent] to [one thing] but, my God, you didn’t know that you were ... going to be taken elsewhere and what was going to happen. So it’s where you’re going.* (Man survivor)

Clinicians are urged to ensure that they are thoroughly familiar with all appropriate sources of information about informed consent.

In many instances, it is necessary to seek consent for each separate component of an examination or procedure:

*Ongoing [consent is required] – it’s not a blanket consent when you’re touching me.* (Man survivor)

The nature of that consent, however, can vary by circumstance. Some men indicated that inquiring about their comfort was synonymous with asking for consent once an examination had begun. For yet other survivors, once trust had been established, the need for the practitioner to repeatedly seek consent for each segment of a procedure was unnecessary:

*As the trust builds in our relationship you would get to a point that perhaps you wouldn’t need to ask me and perhaps I would get to a point where I would say, “You don’t need to ask me for permission any more. We’re now at a point where I trust you and I know you’re not going to hurt me ... But there are some instances where health practitioners need to ask for permission to go to those places.* (Man survivor)

The inclusion of other individuals in any examination/procedure requires additional consent. It is important to inquire about student participation when the student is not present. In addition, even if clients have previously agreed to student involvement for other procedures, always reconfirm their willingness to have the student present, especially during sensitive examinations. Some individuals may agree to have a student present for part of the examination but not for all of it:
rechecking. It doesn’t have to be every minute but perhaps as they move on to another sensitive stage of whatever they’re doing, just recheck. “How are you doing? I’m going to be doing something different now, if at any point you feel uncomfortable, let me know and we’ll proceed from there.” (Man survivor)

To establish a context for ongoing informed consent, practitioners must:

- Allow ample time for patients to explore concerns, ask questions, and decide whether or not they want to proceed;
- Seek consent for each component of an examination or treatment;
- Obtain consent before bringing in students to observe or work with individuals;
- Keep in mind that the onus of ensuring that the client’s consent is truly ongoing is on the clinician;
- Respond to the client’s verbal and nonverbal communication when discerning consent.

6.7 Touch

I rarely go to the doctor; I only go when I absolutely have to and also for physicals. I would just disassociate, that was my only way of coping with that ... any time they touched me. (Woman survivor)

Most survivors we spoke with told us that all forms of physical touch can provoke anxiety. This includes touch from all types of practitioners and in a wide variety of situations, from having blood pressure taken or blood drawn to undergoing a complete physical exam:

Touch is difficult, but if I feel safe, then I can tolerate more. (Man survivor)

Survivor participants agreed that having information before and while being touched is crucial to their sense of safety:
Information and knowing just before you’re going to be touched that it is coming [helps], so, it isn’t sort of a shock to you that you are being touched. (Man survivor)

An understanding of the dynamics of abuse and some of the difficulties behind survivors’ experience of touch can help health care practitioners develop strategies to use during interventions that require touch:

When people used to touch me, it took me right back to the sexual abuse and the physical abuse ... Health care practitioners, if they see their patients sort of backing off or shutting down a bit, they should investigate why. (Man survivor)

For some survivors, the use of touch by the clinician to explain a physical problem or as a component of treatment may be difficult:

[Some clinicians] have automatically assumed that it’s okay for them to go, “Okay, well we’re going to work ... on these muscles [as they touch me] ... because we need to strengthen this because this does this and –” ... It’s not meant [to be sexual touching] ... Once I’ve gone home and calmed down, [and I] thought that they did [not do] anything inappropriately sexual. But at the time, when you first get triggered, it’s an extremely difficult situation to deal with. It triggers a lot of memories ... and then you completely lose whatever you are there for. (Man survivor)

Other survivors spoke of continuously monitoring and reacting to the intent and quality of touch they receive.

If you’re with [health care workers who are] aggressive ... in the way that they touch you, then you’re instantly intimidated and then it’s not safe any more. (Woman survivor)

No one approach to touch is appropriate for every client. Gentle touch, which may be appropriate for some, may be experienced by others as sexually suggestive:

As far as gentle touch goes, you would want a firm but not aggressive touch. Something that’s too soft can be seen as an advance on you. (Man survivor)

Despite the fear and anxiety that many survivor participants experience with touch, some told us about its positive aspects:

I think that touch for healing ... has its place based on my own experience and it helps for rebuilding trust. (Man survivor)

Because touch is such a fundamental issue for survivors of childhood sexual abuse, health care practitioners – before and during any encounters which involve touch – must:

- Recognize that, for many survivors of childhood sexual abuse, no touch is routine;
- Provide patients with information about the reason for and nature of the touch which is involved in any examination or procedure;
- Be sensitive to the intent and nature of all touch, and discuss patient reactions to different types of touch;
- Create a context wherein responses to touch can be freely articulated and the healing nature of touch can be explored.

6.8 Pelvic, breast, genital, and rectal examinations and procedures

Understandably, pelvic and breast exams for women and genital and rectal exams for men and women were cited as being the most difficult parts of a physical exam. Some survivor participants described how these examinations triggered flashbacks for them:

It can trigger ... physical night sweats and severe rectal pain, enormous inexplicable attacks of anxiety. (Man survivor)
Others are unable to tolerate such exams at all:

*I don’t think I would allow anybody to touch me now ... nobody would get an internal on me. No. I will not allow myself to be that vulnerable again.* (Woman survivor)

Participants suggested that practitioners begin by describing the usual sequence of an examination and ask individuals whether they need to adapt it in any way:

*And you know, they take your blood pressure and I said, “You know you’d better do the Pap smear first and then take the blood pressure because right now it will be off the charts and after it will be okay.”* (Woman survivor)

*I think being upright about the actual procedure also made it very difficult for me to talk in general about the other parts of the examination or the other questions they had to ask about the pregnancy or about that kind of thing ... The question-and-answer thing was always before the actual physical exam and I would be really stressed out and really kind of paralyzed feeling, and so I don’t think I ever gave really great information because of that. So maybe having the physical exam first would have helped, get it all over with first, get yourself all back together again ... Or have someone come with you, which I never did, but I suppose that would have helped.* (Woman survivor)

Because there is no single approach that is appropriate in every situation, it is important to:

- Use task-specific inquiry before the exam to learn about anticipated difficulties and negotiate with the individual to minimize discomfort;
- Minimize the time a patient must remain in a subordinate position;
- Drape parts of the body not being examined;
- Allow patients to wear clothing on parts of their body not involved in the examination (e.g., chest, arms, feet, etc.);
- Offer clients a mirror with which to watch the examination or treatment;
- In some cases, suggest to a tense patient that she insert the speculum herself, allowing her to have some control over the intrusion;
- When possible, conduct pelvic examinations with the woman’s head and upper body slightly elevated, as described below:

*There is no single approach that is appropriate in every situation.*

I had been seeing [my family physician] for one and a half years. I kept postponing my physical and the MD noticed that. She kept bringing it up and reminding me until I finally told her that I was frightened of laying flat on my back in a paper gown. She told me that it would not be a problem for me to be partially sitting up throughout the whole examination including the pelvic exam. Now she tells all of her patients that that is an option. She told me that it had been an important conversation for her. (Woman survivor)

### 6.9 Body position and proximity

Both women and men survivors spoke about the difficulty they experienced being in certain positions, partially clad, with a fully clothed clinician standing over them. Health care practitioners can approach the topic of positioning in the same manner as they do other
A time of monumental change for women – a time when the past, present, and future all come together, a time of openness, a time of vulnerability. Being pregnant causes memories of one’s own childhood to surface. Past events are stirred up. The present evokes the paradox of excitement over the baby on the one hand, and fears and anxiety on the other.

According to these authors, some survivors welcome pregnancy as a sign that they are “normal” and develop a growing trust and confidence in themselves as their bodies change to support a new life. For others, however, the experience of pregnancy stirs up memories of past childhood sexual abuse. In her personal account, Christine, an incest survivor, describes not having conscious memory of her abuse until sometime after her third child was born. In retrospect, that knowledge has helped her to understand the difficulties she had with each of her pregnancies: the tears she shed for no apparent reason after every prenatal visit; her severe nausea and vomiting; her long, slow, overdue labours; and the serious postpartum depression. It also explained her life-long shame and distrust of her body, her high need for control, and her life-long struggles with depression. Although she remembers the practitioners who attended her during and after her pregnancies as being caring individuals, none of them ever asked whether she had a history of abuse.

There is some evidence that, compared with women who do not have histories of childhood sexual abuse, survivors: are more reluctant to address their health care needs, have poorer relationships with caregivers, have more anxiety and fear about labour and delivery, report disappointing birthing experiences, are (re)traumatized by the birth experience itself, have more emotional problems in the postpartum period, and experience more problems with breastfeeding and parenting.
I need you to understand this and I need you to help me if I zone out. (Woman survivor)

A practising midwife who participated in our second study told us that she may spend the first two to three prenatal visits just chatting with clients and waits for them to let her know when they are ready for a physical assessment. She tries to minimize invasive procedures and possible triggers as much as possible, performs only necessary interventions, and tries to be flexible (e.g., allowing women to perform certain procedures – such as swabs – themselves).

The principles and guidelines for Sensitive Practice presented in this *Handbook* are a useful foundation for perinatal care. We encourage health care practitioners who work closely with women through pregnancy, birthing, and the postpartum period to read Simpkin and Klaus’s *When Survivors Give Birth* for more specific and detailed guidance concerning the provision of safe and respectful care.

Recommended Readings and Resources – Pregnancy, labour, and postpartum

6.11 Oral and facial health care

Although the guidelines for Sensitive Practice are pertinent to all types of practitioners, there are special concerns for practitioners who work with the mouth, jaw, and face. Because childhood sexual abuse may be oral in nature, many survivors have difficulty tolerating various aspects of oral or facial health care (e.g., the body position they must assume during treatment, the physical proximity of the clinician, and the smells and textures of certain materials such as latex gloves or alcohol):

Too many things in my mouth at once ... You’re making me hold my mouth open too long because you have to do that when somebody’s forcing you to do oral sex, like when you’re a
This means that knowledge of a patient’s past history of abuse is extremely relevant to oral and facial health care:

**Because I was anticipating [difficulty with] a certain procedure I said, “I think you should also know that I am a sexual abuse survivor and ... maybe that’s part of my reaction here.”**

... So if I happen to freak out when he’s poking around in my mouth, that he would have more information there and would know more of what he’s dealing with. (Man survivor)

**Section 7.6 – Triggers and dissociation**

Oral health care can regenerate the feelings of powerlessness and vulnerability that survivors felt as children:

**You have no control because you’re in the chair, your mouth is frozen and you’re pretty much at the mercy of that person.** (Man survivor)

The practitioner who helps the survivor to feel some sense of control during treatment can allay abuse-related fears and increase the likelihood of greater cooperation during treatment. As noted earlier, sharing information and asking permission before performing a procedure can reduce the patient’s feelings of anxiety and powerlessness:

**[The dentist] would talk his way through what he was doing. He would say, “Now I’m going to clean your teeth” or “Now I’m going to spray a little water on that tooth, it may be a bit sensitive.” He doesn’t overdo it but he explains everything he does, so that I have a very clear sense of where he’s going and what he’s doing.**

Many survivors have great difficulty tolerating various aspects of oral or facial health care.

Although practitioners may perceive these frequent explanations and step-by-step consent as repetitious, they are valuable to the apprehensive patient. Intermittently inquiring about a patient’s comfort and following up on negative body language are also helpful, as are establishing hand signals to indicate the need to stop.

**Most of the time [the dentist says] “You know the signals, right?” And I go, “Yeah.” And he’d always review the signals ... “This is what you can do for yes, this is no, this is stop.”** (Man survivor)

Allowing breaks during an appointment or, when possible, breaking a long appointment into two shorter ones can be helpful options for many patients.

**[When I told my dentist that I was having problems that day, he responded,] “Well, what do I need to do? Are you comfortable in the chair? Are we going to need more breaks today?” ... There’s just an unbelievable level of respect with this man. He’s fabulous.** (Woman survivor)

**Section 6.3 – Task-specific inquiry**

Childhood sexual abuse survivors also can feel uncomfortable being in a reclining position in a dental chair with practitioner in close proximity:

**I feel really trapped in the chair, in a very vulnerable position – you know, where you have your mouth open, you’re laid back. For me a lot of my trauma occurred, like, when I was in a laid back position ... and [so having a health care provider] ... over the top of me, I find that very threatening.** (Man survivor)

While supine positioning cannot be avoided, the initial exchange of information and obtaining of
they expect to be judged harshly. This makes it critical that oral health practitioners use a supportive, nonjudgmental tone when presenting treatment options. Instead of reprimanding their patients, oral health practitioners can engender trust by asking how they can best help patients take better care of their teeth.

Survivors indicated that, while oral health care was difficult for many of them, working with practitioners to address the difficulties together often resulted in a positive experience:

*He ... doesn’t ignore what I tell him. He has compassion ... He listened to me. He addressed my situation ... When it’s over ... I feel really great. I really do. Like he’s so gentle, he’s so kind, soft-spoken, yeah, he’s amazing.* (Woman survivor)

To minimize the strain which many childhood sexual abuse survivors experience with oral and facial care, practitioners are encouraged to:

- Undertake the initial exchange of information and obtaining of consent while patients are still sitting upright;
- Opt for gloves made of vinyl or other materials for patients who are anxious about the smell or sensation of latex;
- Establish and use hand signals for “stop” and always respond promptly to them;
- Share information with patients on an ongoing basis;
- Address patient comfort on an ongoing basis by frequently checking in with patients and using task-specific inquiry;
- Follow up on negative body language;
- Allow breaks during an appointment or divide long appointments into two shorter ones;
- Offer patients the opportunity to watch part of the treatment using a mirror;
• Address issues of oral health care neglect in a supportive, nonjudgmental tone and offer to collaborate with patients in finding a way to take better care of their teeth.

Section 6.3 – Task-specific inquiry

6.12 Care within the correctional system

The proportion of childhood sexual abuse survivors is higher within the correctional system than in the general population. Prevalence rates of childhood sexual abuse among incarcerated women range from 47% to 90%;97,152,162 among men, these rates are 40% to 59% (for sexual and/or physical abuse).91,131

Two of the men who participated in our studies were incarcerated in federal institutions at the time of their interviews. They spoke about feeling unsafe when seeking health care not only because of the lack of privacy and confidentiality but also because of standard procedures used within the prison health care system. While the need for strict health care protocols is understandable, it is likely that such protocols dramatically decrease the likelihood that childhood sexual abuse survivors will seek health care:

[When I went to the clinician for a topical nonprescription medication for haemorrhoids, she] wanted to physically check ... I was thinking, “No, no that’s not necessary” ... So she refused me treatment ... [and told me,] “If you’re refusing [to let me examine you] then I can’t give you anything, so I’ll just assume that there’s nothing wrong with you.” (Man survivor)

It is likely that similar difficulties exist for women survivors as well, as has been documented by Pamela Dole,51 a physician who worked in the US correctional system. Although further study is warranted, we urge clinicians who work in the correctional system to examine current practices and seek ways to introduce the principles and guidelines of Sensitive Practice.

6.13 After any physical examination

When ending any interaction, it is essential that practitioners establish a sense of equality with their patients. This will need to be handled differently in different settings:

• In clinic or office settings, see patients when they are fully dressed for health teaching and before they leave an appointment.
• In hospital settings, allow patients who are remaining in hospital garb to regain composure and experience themselves being recognized as a whole person (e.g., shake hands and say good-bye).

• In all settings, invite final questions and, when appropriate, provide a briefly stated plan for future meetings.

6.14 Questions for reflection

• Might any of my current practices be interpreted as insensitive by survivors? What needs to change?
• In what ways might I adapt my own practice to incorporate specific guidelines?
• Do any of these guidelines seem unrealistic or unworkable in my practice? What are some alternate ways of following such guidelines?
• How committed am I to incorporating these guidelines into my routine practice and into the routine practice of those who assist me in my work? What does this level of commitment mean to my clients?
- How aware am I of nonverbal communication of discomfort? Do I follow up on these indicators with my clients?

- Am I aware of resources in my community to which I can refer survivors for care outside my scope of practice? Is this information readily available?
7 Guidelines for Sensitive Practice: Problems in Encounters

7.1 Pain

Pain is a complex issue involving the dynamic interaction of biological, psychological, and social factors that is only partially understood. Research has repeatedly found an association between childhood sexual and physical abuse and increased risk of chronic pain syndromes. An individual may experience pain associated with body (somatic) memories of past abuse in addition to the pain of the disease, illness, or injury for which they seek treatment. 

Research has repeatedly found an association between childhood sexual and physical abuse and increased risk of chronic pain syndromes.

Because an individual’s experience of pain is real, whether or not the pain is consistent with objective findings, it is the clinician’s responsibility to assess the client’s pain in a systematic, thorough, and nonjudgmental manner. The clinician can also:

- Include other practitioners on the treatment team (e.g., mental health practitioners, pain specialists, or pharmacists) to ensure a comprehensive treatment regime;
- Initiate a discussion of other options, including referral to other practitioners who specialize in the management of chronic pain (considering both traditional and complementary practitioners) if an individual’s experience of pain does not remit despite the practitioner’s best efforts;
- Offer a referral, where appropriate, for psychotherapy, clearly explaining the reasons for the suggestion and carefully documenting the details of the discussion;
- Follow up on any referral in future interactions with the client.

7.2 Disconnection from the body

Judith Herman emphasizes the importance of reconnecting with the body in healing from trauma. Being out of touch with one’s body can
make looking after one’s body difficult for a survivor. Indeed, for many survivors, the body becomes nothing more than “a vehicle to get around [in]” (Woman survivor). Such individuals often remain unaware of the messages that their bodies are sending and fail to recognize or attend to signs and symptoms of things such as stress, anxiety, fatigue, or overexertion. These individuals may require specific guidance about activities of daily living and leisure time, physical activity (from doing laundry to gardening), or exercise (either therapeutic or physical fitness training).

For many survivors, assistance from a health care provider to help them become more aware of their bodies may be a critical step in their process of recovery:

“One part of treatment] has been for me to start to get in touch with my body ... I think that a physiotherapist can really affect that [by giving] that supportive invitation to ... come back into [one’s] own body. (Woman survivor) ^143p,256

I needed ... my [massage therapist to] introduce me to my body ... [to] talk to me about my body because I’m not in touch with it. (Man survivor)

Accordingly, health care providers who encounter patients who seem out of touch with their bodies should:

- Repeatedly invite those individuals to focus on their bodies.
- Offer ongoing health teaching about the importance of paying attention to somatic signs and symptoms.
- Provide detailed verbal and written specific instructions for activities of daily living that are problematic as well as for leisure time physical activity. These instructions should include a description of what the activity should feel like and give upper and lower limits for the performance of the activity (e.g., “If your pain increases after making one bed, rest before continuing” or “If you are out of breath, you are doing it too vigorously”).

Feeling out of touch with their bodies can make self-care difficult for childhood sexual abuse survivors.

- Monitor performance and progress.
- Help clients set small, achievable goals to develop neuromuscular skills and understand how to perform the activities correctly.
- Provide careful instructions to facilitate adherence to the treatment program.
- Teach the signs and symptoms of overuse so that survivors can learn how to monitor activity both during treatment and, later, independently of the clinician.
- Suggest a range of strategies to aid self-awareness and connection to the body including: (a) physical activity; (b) somatic-based re-education strategies (e.g., guided visualizations, relaxation exercises, breathing exercises, or yoga); or (c) referrals to other health care providers including complementary health care practitioners.

7.3 Non-adherence to treatment

As stated previously, the inability or apparent unwillingness of clients to adhere to treatment may be related to childhood sexual abuse. Factors such as depression and negative self-perception can lead to unsuccessful courses of treatment for patients and frustration for practitioners. In some instances, the difficulties which survivors experience are directly related to the specifics of past abuse:

There [were] some of the exercises ... that they wanted me to do [after a total hip replacement] ... and one of them that I still

Guidelines for Sensitive Practice: Problems in Encounters
undermines their motivation and sense of agency.

- Where adherence to treatment is particularly important (e.g., for postoperative mobilization), work with individuals to achieve small and reasonable goals (e.g., by ensuring adequate analgesia, teaching splinting techniques, etc.), and acknowledge all successes.

- Remember that blame and guilt are more likely to lead to withdrawal than adherence.

- At the beginning of the meeting, check with clients about reactions during or after the previous meetings, address any problems that have occurred, and answer questions.

- Encourage the view that actively taking care of oneself fosters autonomy and independence.

### 7.4 Appointment cancellations

For many survivors, “walking through the door [for a health care appointment] is a big deal,” (Man survivor) and they cancel appointments as a means of avoidance:

> My wife had been bugging me for a while now, “The dentist has been calling you. You’ve got to go.” “Okay I’ll call her back,” and I don’t call her back. But then eventually ... the adult part of me says, okay you need to go to the dentist ... but the emotional side of me [says] no way I’m going there at all. (Man survivor)

Certainly, cancellations are problematic in that they waste valuable health care resources and are a liability to fee-for-service practitioners and organizations. Nonetheless, it is helpful for service providers to understand why survivors may cancel appointments and, wherever possible, to make changes in their practice environment to facilitate feelings of safety.
changes in their practice environment to facilitate feelings of safety.

To help minimize cancellations, practitioners could:

- Offer “same-day” appointments that would allow survivors to book appointments on days when they feel able to cope. (This can be particularly helpful for oral health practitioners.)
- Work with clients who have identified their apprehension and tendency to cancel appointments to develop a strategy that will assist them.

7.5 “SAVE the Situation”: A general approach for responding to difficult interactions with patients

All health care practitioners encounter difficult situations in the course of their day-to-day practice (e.g., informing a family of the death of a loved one, diagnosing a life-threatening disease or condition, or encountering someone who is angry, anxious, or extremely distressed). These emotionally charged situations may leave practitioners feeling unsure about how to respond. In such instances, reference to the “SAVE the Situation” model may be helpful. The model uses “SAVE” as an acronym for the following four steps: Stop, Appreciate, Validate and Explore. A particular benefit of the “SAVE the Situation” approach is that it can be effective in any difficult situation and is not reserved exclusively for work with survivors.

### TABLE 3

**SAVE the situation**

The acronym **SAVE** is a guide for responding effectively and compassionately in a variety of emotionally charged situations.

<table>
<thead>
<tr>
<th><strong>STOP</strong></th>
<th>Stop what you are doing and focus your full attention to the present situation.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>APPRECIATE</strong></td>
<td>Try to appreciate and understand the person’s situation by using the helping skills of empathy and immediacy. Empathy involves imagining the other person’s experience (thoughts, feelings, body sensations) and communicating an understanding of that experience. Immediacy is verbalizing one’s observations and responses in the moment, using present tense language. For example, ‘Your fists are clenched and you look angry. What is happening for you?’ or ‘You seem upset’ or ‘I doubt there is anything that I can say that will make this easier. Is it okay with you if I sit here with you for a few minutes? If the patient is unable or unwilling to answer, the practitioner can shift the focus to determining possible ways to be helpful (e.g., “How can I help you?”).</td>
</tr>
<tr>
<td><strong>VALIDATE</strong></td>
<td>Validate the other person’s experience. For example, “Given what you have just told me, it makes sense that you feel angry.”</td>
</tr>
<tr>
<td><strong>EXPLORE</strong></td>
<td>Explore the next step. For example, “Who can I call to come and stay with you?” or “This has been difficult for both of us. I am not sure where to go from here. Can I call you tomorrow to see how you are doing?”</td>
</tr>
</tbody>
</table>
7.6 Triggers and dissociation

A trigger is anything (e.g., a sight, sound, smell, touch, taste or thought) associated with a past negative event that activates a memory, flashback or strong emotion. While the focus of this section is on triggers related to abuse, it is not the only cause of this type of adverse reaction to examination and treatment. The suggestions in this section can be used regardless of the origin of the trigger.

[After] surgery on my arm ... the [clinician] would put my arm in water ... [That was something] that my perpetrators had done,

Because triggers are directly associated with a particular event or events, they are unique to each individual. This explains why different stimuli will trigger different people and why a practitioner can never remove or avoid every potential trigger in a practice setting. At the same time, common themes in triggers (see Table 4) are apparent and practitioners are encouraged to consider whether some of these potentially triggering situations can be anticipated. If a patient is able to identify a trigger, the clinician

Differen stimuli will trigger different people and a practitioner can never remove or avoid every potential trigger in a practice setting.

**TABLE 4**

Common triggers

<table>
<thead>
<tr>
<th>Sense</th>
<th>Trigger</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sight</td>
<td>• An individual who resembles the abuser or who has similar traits or objects (e.g., clothing, colouring, mannerisms).&lt;br&gt;• A situation where someone else is being threatened or abused (e.g., a scowl, a raised hand, actual physical abuse).&lt;br&gt;• The sight of an object that was part of the abuse or similar to such an object (e.g., a belt, rope, sex toys) or that is associated with the site where the abuse took place (e.g., a dark room, a locked door).</td>
</tr>
<tr>
<td>Sound</td>
<td>• Sounds associated with anger (e.g., raised voices, arguments, loud noises, objects breaking).&lt;br&gt;• Sounds associated with pain or fear (e.g., sobbing, whimpering, screaming).&lt;br&gt;• A situation in which the survivor is being reprimanded.&lt;br&gt;• Sounds associated with the place or situation before, during, or after the abuse occurred (e.g., footsteps, a door being locked, a certain piece of music, sirens, birds chirping, a car door closing).&lt;br&gt;• Anything that resembles sounds that the abuser made (e.g., particular words, phrases or tone of voice, whistling, cursing, groaning).</td>
</tr>
<tr>
<td>Smell</td>
<td>• Odours associated with the abuser(s) (e.g., cologne or after-shave, tobacco, alcohol, drugs).&lt;br&gt;• Odours associated with the place or situation where the abuse occurred (e.g., mildew, petroleum products, food odours, outdoor smells).</td>
</tr>
<tr>
<td>Touch</td>
<td>• Any type of physical contact or proximity that resembles the abuse (e.g., touch on certain parts of the body, touch that comes without warning, standing too close, the sensation of breath on the skin, the manner in which someone approaches).&lt;br&gt;• The sensation of any type of object that was used during abuse (e.g., ice, gel similar to lubricant or semen, the sensation of equipment that is reminiscent of restraints used during abuse).</td>
</tr>
<tr>
<td>Taste</td>
<td>• Any taste related to the abuse (e.g., certain foods, alcohol, tobacco).</td>
</tr>
</tbody>
</table>
and patient can problem-solve together to either avoid or minimize that trigger during future interactions.

Clinical practice incorporates many experiences in addition to touch that may trigger a negative response in a survivor even though they seem innocuous to the clinician. Survivors described triggers such as the use of water, ice, traction, or ultrasound gel. They also spoke about medical procedures and treatments during which they had to remain immobile or silent or heard others crying out with pain or anxiety, reminding them of abuse experiences. Other participants told us that a practitioner’s body language or reprimands for behaviours interpreted as deliberate non-adherence to recommendations could also trigger intensely negative experiences.

Survivors may or may not be aware of their triggers and may realize that they have been triggered only after they have had this experience. Individuals may also be triggered whether or not they have conscious memory of past abuse or have disclosed to anyone:

[During] my first experience [with this type of practitioner], they didn’t have any Kleenex, and the minute [the clinicians started] touching me I just started sobbing, without having any idea of ... why. (Woman survivor)

When he did the physical examination I just basically dissociated myself from my body and I never had any idea why ... or how I did it. But looking back now, I used to do that quite a bit. After the examination was over I had no idea what he said to me. The only thing I wanted to do was get out of there. I felt extremely violated. (Man survivor)

Survivors stressed that it is important that all clinicians have a general understanding about triggers and how to respond to an individual who is triggered:

The flashbacks that could happen while you’re having an exam. The not being present in the moment ... It would be helpful for a [clinician] to be able to help bring a patient back into the present moment and give them the time to sort through what’s going on in their head. (Woman survivor)

If you have a guy crying in front of you and especially if he’s a victim, [if you understand triggers], at least you can have some type of understanding of where this person’s coming from. (Man survivor)

Health care practitioners should be attuned to the following behaviours, which may be nonverbal indicators of discomfort, distress, or dissociation:

- Rapid heart rate and breathing (breath holding or sudden change in breathing pattern may also be seen);
- Sudden flooding of strong emotions (e.g., anger, sadness, fear, etc.);
- Pallor or flushing;
- Sweating;
- Muscle stiffness, muscle tension, and inability to relax;
- Cringing, flinching, or pulling away;
- Trembling or shaking;
- Startle response.

These behaviours are probably best understood as “freeze-fight-or-flight” responses to the perception of a threat (i.e., sympathetic nervous system arousal).

The following responses may be clearer indications of dissociation:

- Staring vacantly into the distance;
- Spacing out or being uninvolved in the present;
- Being unable to focus, concentrate, or respond to instructions;
- Being unable to speak.

After being triggered into a dissociative state, an individual may seem confused or vague and ask questions such as “Where was I?”, “What did I just say?”, or “What just happened?” However,
it is possible that the clinician and even the clients themselves may not know that they have dissociated. Indeed, some survivors only discover as adults that they dissociate under stressful circumstances:

*The health care practitioner would come into my personal space and... I would just dissociate. She'd touch me and then I'd just be gone. She worked with a lot of women who were survivors and she knew it. She'd just stop and say, “Where did you go?” And I didn’t have a clue what she was talking about. But over the years I started getting a clue.* (Man survivor)

Now, *clinicians* don’t have to handle the *whole* crisis, but they do need to know how to recognize [it]. And how to make a referral in a nice way [by saying, for example] “Do you see your counsellor tomorrow?” or “Is there someone you can talk to?” ... They wouldn’t need to go beyond [their scope of practice], but [it is helpful] if they can recognize what can happen when a woman is going through a flashback ... [and know] how to ground a person. It’s not hard; ... [it’s] just basic humanity and reassurance. You know, “You’re okay, it’s safe here,” or [validating] the energy and the courage that it takes to go through [the specific intervention] ... And [they can say,] “Yes, [this treatment] can trigger memories, and it can be really disturbing and distressful, and what you’re feeling is normal.” (Woman survivor)

To support clients who have been triggered and ensure that they do not leave the encounter feeling disoriented or embarrassed about their reactions to treatment, practitioners should:

- Follow the SAVE protocol;
- Orient clients to the present by reminding them where they are and what was happening when they began to have trouble staying present;
- Encourage slow, rhythmic “4-6 breathing” (inhale to the count of four and exhale to the count of six) and (if possible) sitting up and placing their feet on the floor;
- Remind individuals to keep their eyes open and to look around the room;
- Encourage patients to notice physical sensations (e.g., the feeling of their back on the chair and their feet touching the floor, or the sensation of the air on their face).

As clients become more oriented and responsive:

- Do not touch them;
- Offer verbal reassurance in a calm voice;
- Avoid asking complicated questions or giving complex instructions; instead, ask simple questions to try to connect with the person (e.g., “Are you with me?”, “Are you following me?”, “Do you have ways of staying present?”);
- Offer them a glass of water;
- Allow them the necessary time and space to regain their equilibrium (a quiet room may be helpful);
- Normalize the experience. If the patient has disclosed abuse prior to this incident, let her or him know that health care interventions commonly trigger flashbacks or emotional responses, but do not ask for details of past abuse that may have contributed to being triggered. If the patient has not disclosed abuse, frame the normalizing comments in terms of anxiety that many people feel when seeing health care practitioners;
- Ask what the clients need right now (e.g., do they want your company, or would they rather be left alone);
- Offer continuity of care (i.e., if time constraints prevent you from staying with upset clients as long as you would like, explain this and ask if someone else can help, such as another staff member or a friend whom you could call).
Being triggered can be a frightening or bewildering experience. Some clients may benefit from talking about the experience with someone. Thus clinicians should:

- Inquire about whether the patient has someone to offer support and whether they would like to contact that person now (e.g., “A new exam like the one we were doing today can be scary for many people and can bring about very strong emotions, as you just experienced. Sometimes it helps to talk about what happened. Do you have anyone you can process this with? Would you like to call this person to be with you now?”).

- Find out whether patients would like to explore what has happened; if they have no one to talk with, ask them whether they want a referral to a counsellor or other community resource and whether they know about telephone help lines that exist in your community.

- Ask whether the client feels able to continue the examination or treatment.

A person who has been triggered or has dissociated may not retain or recall important information shared by the clinician. Thus, it is helpful for practitioners to:

- Repeat all instructions;

- Write down instructions and recommendations in clear language.

For individuals who have repeated experiences of dissociation during their interactions with clinicians:

- Suggest that they use a notebook to write information, instructions, and suggestions;

- Share with clients the responsibility for ensuring that essential information is recorded before the end of the interaction.

The next time the practitioner sees the client who has been triggered or dissociated:

- Discuss the experience with clients to ensure that they are feeling better and to reaffirm the message that the event does not alter the esteem in which they are held;

- Problem-solve with clients to identify what to avoid or modify in the future to prevent further triggering, keeping in mind that they may or may not be able to identify the trigger of a particular incident;

- Learn from the individuals what techniques they use to stay present and grounded, including any reminders or instructions that you can give them;

- Suggest – if the severity of the client’s reactions and subsequent difficulty so indicates – a consultation with a mental health practitioner to develop additional strategies for coping with triggers.

Some survivor participants suggested that practitioners offer general cautionary messages to clients about adverse reactions to procedures or treatment that are invasive or uncomfortable (e.g., pelvic and rectal exams and dental work):

_Something that my orthodontist may have never realized, for me with that history of abuse, [is that] when I got the braces on, for three nights in a row I just had horrible nightmares. I was phoning my counsellor and saying, “Can I book an appointment, get in right away?” Because I didn’t have a clue what was going on … All of a sudden I’m having nightmares being that little kid again because of all this prodding and pulling going on in my mouth. I would want an orthodontist handing out a leaflet going, if you’ve had sexual abuse, keep in mind this could give you nightmares or this could trigger you._ (Man survivor)
If health care providers are shaken or upset by the triggering or dissociation of a client, they should talk with a colleague, a supervisor, or someone within their support system. This can be done without breaching confidentiality.

Section 3.5 – Specific behaviours and feelings arising during health care encounters
Section 5.9 – Practitioners’ self care

7.7 Anger or agitation

Anger is my initial response to almost everything ... I try to hold on to myself, which I do much better than before I was 40. But initially my reaction is to get angry. (Man survivor)

Many men survivors and a few women survivors talked about responding with anger when they are anxious or fearful or have been triggered:

Anger shows up often when you are triggered – like [when] somebody touches you in the wrong place. (Man survivor)

While it is generally easy to respond compassionately to someone who is sad or afraid, anger often elicits the opposite response – defensiveness, irritation, or withdrawal. This type of response, however, can leave survivors in an even more difficult situation:

You are frightened and everybody is frightened of you. (Man survivor)

Health care providers will benefit from recognizing the connection between anger and past abuse for some survivors as well as from realizing that anger (an emotion) and violence (a behaviour) are distinct entities, not to be confused or seen as one response. Participants recommended that practitioners use the SAVE guidelines to understand the cause of the anger. They advised against trying to control agitated patients’ behaviours; rather, they suggested that the clinician: (a) allow clients time to cool down; (b) reflect their observations back to the clients; and (c) work with them in seeking a solution to the problem (i.e., “Don’t dictate, negotiate!” (Man survivor)).

A situation can quickly escalate if a practitioner responds to an angry or agitated client with defensiveness or anger. Managing one’s own anger is critical to interpersonal effectiveness. Although many institutions and organizations have established policies to deal with angry and violent patients, it remains crucial that health care providers:

- Manage their own feelings of anger;
- Pay attention to personal safety (e.g., do not stand too close, do not make quick or sudden movements, identify an escape route);
- Adopt non threatening body language (e.g., stand with arms uncrossed, at a slight angle to the person to avoid the experience of face-to-face confrontation);
- Speak slowly in a low voice, breathe slowly and rhythmically.
- Encourage agitated individuals to relax and assure them that you are interested both in listening to their concerns and in helping them find solutions to their problems.

Anger often elicits responses of defensiveness, irritation, or withdrawal.
8.1 The challenge of disclosure for survivors

Choosing whether to disclose depends on where you are in your journey. Because sometimes in your journey you don’t want them [health care providers] to know [you’re a survivor]. (Woman survivor)

Although this discussion speaks of survivors collectively, survivors are not a homogenous group; each survivor is a unique individual with a unique history and point of view. While survivors may or may not disclose their histories, their abilities to recall the abuse and their places in the journey towards recovery consistently play significant roles in disclosure. For example, most of the study participants have always had clear memories of the abuse they experienced, while a smaller number only began to remember the abuse in adulthood. Some attempted to deny to both themselves and others that the abuse occurred:

I just buried it and pretended that it didn’t happen ... and sort of just [said to myself], “No, no-how could that affect my life?” And it wasn’t until last year I really started to realize that it did affect my life. I knew it wasn’t right at the time, way back, but I didn’t know that it could potentially have the effect that it had. (Man survivor)

Some survivors also told us that they had always remembered their childhood abuse, but did not identify it as abusive (believing that what happened to them happens to all children) until some new learning prompted them to reconsider their experience:

My awareness of my childhood sexual abuse only dates from about nine or ten years [ago].

[i’m] not saying that I didn’t remember things that happened to me. I did, but I didn’t appreciate the dynamic that was there and I just sort of thought of them as early sexual experiences and said [to myself], “Well, doesn’t that happen to everyone?” Then you suddenly discover that no, it doesn’t happen to everyone. The real dynamic just really clicked one day and it really hit me hard. (Man survivor)

Participants described a number of factors that influenced their decisions about whether to disclose to practitioners, and also how much and what information they shared. Some did so spontaneously early in the relationships, while others held back until they felt more comfortable with the clinicians. Still others chose not to disclose at all.

Survivors who disclosed their abuse spontaneously (i.e., not in response to questioning by a practitioner) did so in the hope that the information would help the practitioners to understand them better.

[i disclosed so that the clinician would] have some of the understandings of the feelings that are associated with that part of the physical exam ... – the shame and the guilt and the things that you have going on inside your head, the flashbacks that could happen while you’re having an exam, the not being present in the moment. (Woman survivor)

For many survivors, disclosure is a process. Unlike survivors who want to “get it over all at once,” others prefer to reveal their history gradually over time, often so that they can take control of the timing and pace of disclosure:

[My doctor’s response] helped me, little by little, disclose more of my deep dark secrets
Finally, some survivors want to avoid having to disclose repeatedly and take a proactive approach to the issue:

At this point in my life I think differently [than I used to about disclosing], I want “survivor” written on the front of my chart so that [clinicians] know and recognize that I want to be treated sensitively. Then, if a new [person] in the practice sees me it would be a reminder to them. Other survivors may not want that, but I think it would be great if I did not have to disclose every time I see a new [practitioner].

(Woman survivor)

A reluctance to disclose may relate to: (a) survivors’ feelings about themselves; (b) pressure from families, friends, or abusers to remain silent; (c) their fear of negative responses; and/or (d) the sense that their practitioners do not have the time to listen or seem unaware of the potential long-term health implications of violence.

Many individuals spoke about how their own feelings of shame and guilt affected their attitudes towards disclosure:

There’s a whole lot of shame [about having been victimized] … and disclosing that.

(Woman survivor)

Others told us of the vulnerability they feel when disclosing:

Every time you disclose, you expose yourself.

(Woman survivor)

Previous experiences with disclosure play a major role in survivors’ decisions about disclosure. Many were reluctant to say anything because they feared a negative reaction, particularly rejection:

I’m really hesitant on mentioning it to people, especially … [to health practitioners] – I don’t want to start talking about it or mention it and get that rejection. ‘Cause that’s the worst. ‘Cause then I clam up and I – my headaches will probably get worse and everything will just get worse.

(Woman survivor)

Others fear being blamed for the abuse or being judged:

One [practitioner] that I saw … reacted with insensitivity, by asking me, “How did you let it happen?” In the moment I felt revictimized and took all of the blame for what happened. That really had an impact on me.

(Woman survivor)

Many male survivors, in particular, are fearful that if they disclose past abuse a clinician will assume that they are also perpetrators:

I called the hospital to talk about sexual abuse and they thought that I was the abuser and referred me to domestic sexual abuse centre.

(Man survivor)

Both men and women whose abuser(s) were women were reluctant to disclose for fear of not being believed:

Female survivors of female-perpetrated abuse … experience disbelief as to the likelihood of having been abused by a woman.

(Woman survivor)

If it was with a woman it’s, “Well aren’t you mistaking it for nurturing?”

(Man survivor)

Finally, both survivor participants and health care practitioner participants identified practitioners’ apparent lack of time as a huge barrier to disclosure:

I was almost 60 when I started [to deal with issues of sexual abuse] and it came to light after a lot of very significant [psychotherapeutic] work of mine … So these are deep things. In other words, this is a deep question and to think of it in terms of a 15-minute segment [with a clinician] is hard.

(Man survivor)

8.2 Possible indicators of past abuse

While there is no single indicator or cluster of symptoms and/or behaviours that provides evidence of past abuse, there is a growing body of research that documents a relationship between
adverse childhood experiences and certain behaviours and/or experiences in later life. Some of these include:

- Avoidance of all health care practitioners and/or health serving agencies;
- Repeated cancellations of appointments;
- Repeated postponement of a physical exam;
- Poor adherence to medical recommendations;
- Chronic unexplained pain (e.g., headache, pelvic, back, muscular);
- Unexplained gastrointestinal symptoms/distress;
- Disordered eating, obesity, or wide fluctuations in weight;
- Sleep disturbances (insomnia, hypersomnia);
- Sexual problems (e.g., avoidance, many sexual partners, unsafe sex practices);
- Alcohol or drug misuse;
- Depression;
- Pattern of difficulty in interpersonal relationships;
- Self-harm behaviours and/or suicide ideations/Attempts;
- Posttraumatic Stress Disorder or other anxiety problems;
- Dissociative states (blanking out, long silences).

Recognizing clusters or patterns of these behaviours and symptoms along with inconsistencies or gaps in information provided by the patient should alert a clinician to consider the possibility of abuse or violence:

But I would ask [practitioners] to go a step further, to [talk] ... to men, particularly males who have addiction problems, who have eating disorders, sleep disorders, depression, anything that has to do with emotion, emotional things or mental health issues. I think it’s important that these [clinicians] ... get trained to be able ... to identify [behaviours that may be related to past abuse] and to be up on what the actual symptoms are. (Man survivor)¹⁵⁹

It is crucial that health care providers be aware that these indicators, although clearly suggestive of abuse or psychological trauma, may actually stem from other causes. Abuse is not always the source of these behaviours; nonetheless, inquiry about a history of childhood sexual abuse is essential.

8.3 Inquiring about past abuse

A growing body of evidence indicates a relationship between abuse or violence and health problems. Our studies further demonstrate a range of ways in which past abuse can negatively affect survivor-practitioner interactions. Accordingly, inquiring about violence and abuse should be an integral part of collecting a health history:

I think it’s important that [health care practitioners] ask questions about abuse as part of a medical history, particularly of women, and I think that anyone dealing with women’s pain who doesn’t ask questions about violence in a woman’s life is not doing their job. I feel that very, very strongly. (Woman survivor)¹⁶⁴p.93

Recognizing clusters or patterns of these behaviours and symptoms along with inconsistencies or gaps in information provided by the patient should alert a clinician to consider the possibility of abuse or violence.
times I will ask if stress or being emotionally upset causes their symptoms to worsen. If they respond “Yes,” then I will ask what the greatest causes of stress are for them (“Is the source of your stress: home life, relationships, work, school, finances, family issues etc.?“). Once the patient confirms that stress is a factor and that they can identify what their main stress reaction triggers are, then I will ask if they have a good support system (“Do you confide in friends, significant others, other family members?”). I next will ask if they actually use their support system. Many patients will respond with comments such as “Not as much as I should” or “Yes, and I think that they are tired of listening to me.” At this point I am able to intervene by explaining to the patient that I have a good referral network, and that perhaps they should consider seeing a counsellor. I reassure the patient that I do not necessarily require any details regarding their stress, but many patients will spontaneously divulge ... In short, history taking allows me to develop a relationship with the patient. During history taking, when the patient feels heard and cared for, then the patient will often disclose childhood sexual abuse. Patients are always reassured that they are in control of everything that takes place during their visit. Communication is established during history taking and is reinforced during examination and treatments.

Inquiring about violence and abuse should be an integral part of a health history.

As one practitioner participant reported:

Most patients present for chiropractic care for pain (lower back pain, neck pain, headaches). During history taking I ask if they can identify aggravating factors. Sometimes patients will relate stress as an aggravating factor. At other times I will ask if stress or being emotionally upset causes their symptoms to worsen. If they respond “Yes,” then I will ask what the greatest causes of stress are for them (“Is the source of your stress: home life, relationships, work, school, finances, family issues etc.?“). Once the patient confirms that stress is a factor and that they can identify what their main stress reaction triggers are, then I will ask if they have a good support system (“Do you confide in friends, significant others, other family members?”). I next will ask if they actually use their support system. Many patients will respond with comments such as “Not as much as I should” or “Yes, and I think that they are tired of listening to me.” At this point I am able to intervene by explaining to the patient that I have a good referral network, and that perhaps they should consider seeing a counsellor. I reassure the patient that I do not necessarily require any details regarding their stress, but many patients will spontaneously divulge ... In short, history taking allows me to develop a relationship with the patient. During history taking, when the patient feels heard and cared for, then the patient will often disclose childhood sexual abuse. Patients are always reassured that they are in control of everything that takes place during their visit. Communication is established during history taking and is reinforced during examination and treatments.

While our own research and that of others makes it clear that health care practitioners have a professional and ethical responsibility to inquire about abuse or violence, it is important to understand this statement in relation to the debate regarding the evidence pertaining to inquiry about/screening for interpersonal violence. It is also important to acknowledge that not all survivors want to be asked about past abuse and may choose not to disclose:

If I wanted to tell him, I’d tell him. It’s not his business. (Man survivor)159

As long as health care providers respect the wishes of survivors who prefer not to disclose a
Both women and men described a number of factors that might encourage disclosure. They look for signals that the clinician has an understanding of the effects of interpersonal violence, including posters and pamphlets (directed at both women and men) prominently displayed in waiting rooms, washrooms, and examination rooms. Survivors also stressed the importance of feeling safe, and trusting their practitioners:

My doctor made me comfortable from the beginning so I felt I had someone to talk to. I've been married for 28 years and I wasn't even able to tell my wife, but I was able to tell him. If I wasn't able to tell him I don't know if I would have been able to move in the direction of recovery. (Man survivor)

There was this one specific [practitioner who] was just so, so kind . . . that person would definitely be someone that I would not have a problem sharing, you know, what had happened to me, what I had experienced. (Woman survivor)

Survivors emphasized the importance of confidentiality in their decision to disclose:

I guess the primary issue is . . . confidentiality. [I need to know,] are you going to tell anybody? Are you going to do anything with the information? (Man survivor)

Appendix I – The Evidence Debate Pertaining to Inquiry about Interpersonal Violence

For many practitioners, the first step towards routine inquiry about interpersonal violence is an attitudinal one. Studies have shown that barriers to inquiring about interpersonal violence include: (a) a lack of knowledge and training about the topic and how to ask relevant questions; (b) lack of privacy and time limitations; (c) the belief that abuse is not a problem for their patients; and (d) frustration with being unable to help the victim. A clinician’s own experience with violence might also factor into an unwillingness to address the topic with patients. Nonetheless, routine inquiries about interpersonal violence are fundamental to Sensitive Practice:

Surely [practitioners] realize that it’s a part of who I am and it needs to be acknowledged, and it does have an impact in terms of how I need to be treated. (Woman survivor)

Guidelines for Sensitive Practice: Disclosure
The therapeutic relationship and the health care environment are crucial factors in the inquiry about past abuse. The Society of Obstetricians and Gynaecologists of Canada's clinical practice guidelines offer a valuable reminder to clinicians about the therapeutic relationship: “Several validated questionnaires exist for enquiring about [interpersonal violence]; however, the nature of the clinician-patient relationship and how questions are asked seem more important than the screening tool.” Regarding the environment, survivor participants emphasized that privacy and clearly visible and available information (e.g., posters or brochures) convey the impression that a practitioner acknowledges the relationship between interpersonal violence and health.

**Verbal inquiry.** There is no one correct way to ask about a history of childhood abuse. Direct approaches are a relief to some survivors, but may be too intrusive for others. Introducing questions in a way that relates past abuse to health and health care provides context and rationale. Practitioners could draw on the following statements as possible lead-ins to an inquiry of childhood sexual abuse history:

- “Research tells us that child sexual abuse among both girls and boys is much more common than was once believed. We also know that it can have long-term health effects.”

- “Is there anything in your history that makes seeing a practitioner or having a physical examination difficult? If there is, I would like to hear about it so that we can work together more easily.”

- “Some women (or men) want to talk with their health care providers about very personal or difficult topics. If you do, I am open to hearing about them.”

Statements and questions such as these may open the door to disclosure, either in the moment or at some later time. If an individual hesitates or seems very reluctant to respond, another effective response from a clinician would be something such as:

- “I know these things can be hard to talk about. I think it is important to ask because there is growing evidence that violence and abuse can affect a person’s health and create difficulties when they see health care practitioners. You don’t have to discuss this with me if you don’t want to. If you do, I can work with you to ensure you are comfortable when you see me and to get whatever support or assistance you need.”

Regardless of how the questions are framed, participants told us that trust in their healthcare provider influenced their decision to disclose:

> I had one [practitioner] ask me, “Was there any trauma in your childhood or lately that could cause these symptoms?” And right then, I thought, “Oh, okay. I can talk about it. And I’m not going to get rejection.” (Woman survivor)

Some men survivors told us that they were unclear whether clinicians were asking if they were victims or perpetrators of sexual abuse when asked questions such as, “Have you encountered sexual abuse?” Therefore, they urged health care practitioners to clearly ask if the man is a victim of past abuse.

Most of the survivors in our research, both women and men, indicated that they did not want to discuss the details of their abuse with their practitioners:

> Some people press for more info upon disclosure and that is invasive and unacceptable. (Woman survivor)

**Written inquiry.** Survivors in our studies varied in their views about the merits of written and oral
When survivors disclose their history of abuse, it is usually because they hope that something positive will come from it. If practitioners do not respond, survivors may interpret the silence as an indication of lack of interest, which may deter them from mentioning it again. Moreover, they may stop seeing that particular clinician or, in the extreme, avoid all health services:

I told the [health care provider] about my history of abuse. She didn’t acknowledge [it] ... She just kept right on going with what she was doing ... Oh boy! If somebody says it, then you’ve got to acknowledge it. Because then what that says to me is that it’s not valid, it’s not important, it doesn’t have anything to do with us. (Woman survivor)164p.95

Express empathy and caring. Survivors also want to know that their practitioners care about them. Simple statements of empathy and concern can convey both compassion and interest:

He just looked at me and he said, you know, I’m really sorry this happened to you. And that was the best thing he could have said. (Woman survivor)159

Communicating to survivors that they have been heard and believed is crucial whenever survivors disclose. While follow-up is also important (as the next section indicates), the practitioner’s immediate verbal and nonverbal responses to disclosure can have a tremendous impact on the survivor.

Accept the information. Individuals need to know that their health care providers have heard them, have accepted the information, and believe that children are never responsible for abuse:

His response was first one of acknowledging what I said and, you know, genuinely looking like he cared and kind of going with that and not really pushing anything, not giving me advice or telling me what to do but, you know, just kind of going slowly with me through that. And I found that was excellent. (Woman survivor)

The most prudent strategy is for health care providers to use both written and verbal forms as part of every health history.

Clarify confidentiality. Confidentiality is a vital concern for many survivors. Although a clinician may have already discussed it previously, following a disclosure of abuse, health care practitioners need to repeat information about the level of confidentiality that they can extend. For example, the clinician might say, “Because you are an adult now, I am under no legal obligation to report this to police or a child welfare agency” and “I think it is important to write something about your

8.4 Responding effectively to disclosure

Well, for one thing, it’s really important [to tell survivors] ... that you believe them, because this might be the first person they’ve told. And also, it’s really important to accept them as a person. You can say whatever your real feelings are. [For example,] “I’m really sad to hear that.” (Woman survivor)143p.258,164p.95

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I remember feeling comforted by her, probably by her words. She probably said, “It’s okay to cry” or she might have even rubbed my arm. I remember her telling me that she was going to give me a phone number where I could call so I could talk to somebody about it, which she did. She handled it very professionally. (Woman survivor)

Clarify confidentiality. Confidentiality is a vital concern for many survivors. Although a clinician may have already discussed it previously, following a disclosure of abuse, health care practitioners need to repeat information about the level of confidentiality that they can extend. For example, the clinician might say, “Because you are an adult now, I am under no legal obligation to report this to police or a child welfare agency” and “I think it is important to write something about your

Guidelines for Sensitive Practice: Disclosure
childhood history in the chart. What would you like me to put down?”

Section 8.7 – Legal and record-keeping issues

The most important thing is, “Whatever you say is confidential with me.” Because confidentiality is so huge. (Man survivor)

Acknowledging the prevalence of abuse. Understandably, many survivors feel very isolated and alone in their experience. Having health care providers demonstrate awareness about the prevalence and long term effects of childhood sexual abuse normalizes the experience for survivors and may reduce their sense of shame. For example, a clinician might say, “We know that as many as one in three women and one in seven men are survivors of childhood sexual abuse. It is sad to realize that so many children have suffered in this way.”

Validating the disclosure. Health care practitioners must validate the courage it took to disclose and communicate that they believe what they have been told. Visible distress needs to be acknowledged (e.g., “I see that this is painful [distressing, disturbing] for you right now. What can I do to help?” or “It is okay if it takes more than one visit to do a complete examination”). Failure to validate the individual’s experience, silence, or judgemental comments can be shaming and contribute to a reticence to disclose in the future:

[It is important] to validate that experience because ... [it is hard] to keep that buried for 20 years and then bring it out and start talking about it and then look across and see a look of what you might perceive to be disbelief in somebody’s eyes and you’re wondering inside yourself, you know, ... maybe I am crazy and it didn’t really happen or it wasn’t like that or, you’re supposed to be a man and it wasn’t that bad and just shake it off and carry on right? (Man survivor)

Address time limitations. Time pressures are one of the biggest impediments to disclosure. If individuals disclose a history of abuse and the health care provider can spend only a few minutes with them afterward, it is important that the time constraints are communicated in a way that will not leave survivors feeling dismissed or that they have done something wrong by disclosing (e.g., “Thank you for telling me about being abused. I can only imagine how difficult things have been for you. I have another patient waiting – do you want to book a longer appointment later this week?”).

Section 4.2 – Second Principle: Taking time
Section 6.5 – Time
Section 8.5 – Additional actions required at the time of disclosure or over time
Section 8.6 – Responses to avoid immediately following a disclosure

Offering reassurance. Because individuals who have disclosed have shared some very personal information, they may feel vulnerable and exposed – both at the time of the disclosure and during future encounters with the practitioner to whom they have disclosed. To minimize this sense of vulnerability, practitioners can reassure survivors that they applaud the courage it takes to talk about past abuse and that the information that has been shared will be useful in providing appropriate health care.

Collaborating to develop an immediate plan for self-care. Some survivor participants identified unsettled feelings or flashbacks of their abuse as an immediate after-effect of disclosure:

I was triggered more, and I was getting more flashbacks after [disclosing the abuse]. (Man survivor)

Accordingly, health care providers should caution individuals who have just disclosed to be prepared for these feelings. They should then work with survivors to make a specific plan for self-care
Section 8.5 – Additional actions required at the time of disclosure or over time

Ask whether this is the patient’s first disclosure. As well as responding to a disclosure as outlined above, health care providers can inquire whether the patient is disclosing for the first time. By asking “Have you talked with anyone else about this?” practitioners can get a sense of whether the survivor has previously taken any steps to address the abuse. An answer of “No, I have never told anyone before today,” as compared to “Well, my counsellor knows and suggested that I tell you,” can help clinicians to shape their next response. It may also help them learn what supports the clients have in place and what they may need.

8.5 Additional actions required at the time of disclosure or over time

Either immediately following the disclosure or during the next interaction, health care providers should seek to understand the survivors’ reasons for disclosing and determine what (if anything) they want from the practitioners. It is also important to clarify the survivors’ general expectations of the clinician and to explore any implications that the disclosure has for the survivors’ health care. Such questions need to be asked in a manner that indicates clear support for the individuals’ choice to disclose and may provide a bridge to discuss ways to maximize their feelings of safety and comfort. While such discussions may take some time (and be spread over a few interactions), the information which comes from them will provide a basis for future interactions:

When I came in, [the clinician] said, “I did some reading up on your condition,” and he said, “This is what we’re going to do.” He says, “We’re going to work out a system, okay, so that I know if you’re having trouble and you need to stop.” (Woman survivor)

In working out this plan, clinicians should encourage individuals to:

- Include activities and coping strategies that have been successful (i.e., are supportive, comforting, or help the individual to manage distressing emotions).
- Be specific and realistic, and include things that are easy to implement in a moment of distress. An unspecific plan (e.g., to take it easy for the next few days) may be too ambiguous to translate into meaningful activity, whereas a more specific plan (e.g., to call a specific support person or engage in a specific activity, such as going to the gym, meditating or praying, writing in a personal journal, or attending a self-help group meeting) gives survivors clear direction.
- Include ideas about what to do if the usual coping strategies do not work. This step is particularly important if the individual has a history of depression or self-harm. It might involve calling a health information line or crisis line or going to the emergency department of the local hospital.

Recognize that action is not always required. Health care practitioners tend to be problem-oriented and may respond to disclosure as a problem that requires immediate action or resolution; however, survivors may simply want the clinician to have the information. Survivors who have just disclosed may not necessarily expect clinicians to do anything except to be present with them in the moment. While it is important to ask survivors if there is anything they want done related to their disclosure, it may be preferable to identify a later time for discussion about what actions (if any) the survivors want from the practitioner.

Survivors who have just disclosed may not necessarily expect clinicians to do anything except to be present with them in the moment.
Practitioners might say, for example, “Knowing this will help me care for you better. Can we talk about things that might make you more comfortable during your appointments?” or “Is there anything I can do differently?” The ensuing discussions may lead to disclosures of task-specific issues as survivors gradually feel freer to express their needs or preferences. As difficulties are identified, clinicians can integrate changes into the individual’s ongoing care. Regardless of what is accomplished, health care practitioners should not assume that all issues have been dealt with in one or two discussions; rather, they should check in with their clients throughout each interaction and make repeated invitations for feedback:

[The practitioner could say,] “Just let me know [what you need]; the lines are open. I know this [abuse] happened and if you need to talk about it or have any questions [you can talk about them with me].” (Man survivor)

[After] I told him I was a survivor ... he always questioned if I was comfortable doing anything ... Communication was more [important]. (Woman survivor)

Some survivors hope for a response that is beyond the clinician’s ability or scope of practice. It is therefore important for practitioners to be clear about what they can and cannot do to help. If clinicians feel that individuals require assistance beyond that which they can offer, then a referral to someone more able or qualified may be suggested.

Most survivors recognized that disclosing their history of abuse was important to both their health and their health care. Nevertheless, many were concerned that, once they had disclosed their history of childhood sexual abuse, their health care practitioners would tend to attribute their health problems to the abuse before thoroughly investigating other possible reasons for the problems:

[Practitioners] should never assume. Just because I was abused, that doesn’t or shouldn’t rule out the possibility that there could be something physical and serious that is wrong. That’s one of the reasons I don’t like to tell ... health practitioners about my abuse. They tend to write everything off as nerves and don’t even check to see if the problem is something else. (Woman survivor)

While an abuse history may contribute to some illnesses, it is the clinician’s responsibility to ensure that health problems are investigated thoroughly for all patients.

Because of the vulnerability that they felt after disclosing their abuse histories, some survivors were wary about being referred to other health care practitioners. Although clinicians typically see referrals as a normal and reasonable action to ensure accurate diagnosis and treatment, survivors may think that the referral implies that their practitioners cannot take care of them because they are “too complicated.” As well, survivors may feel uncomfortable or anxious about having to meet one more clinician, whom they do not yet know or trust.

All patients have the right to make an informed choice about the health care practitioners with whom they will work. Thus, before making referrals, practitioners are encouraged to discuss the issue with their clients in order to come to an agreement on a new practitioner. These discussions may be very significant for survivors who, for example, are uncomfortable working with clinicians who are the same gender as their abuser(s). Whenever possible, practitioners should refer to health care providers who are knowledgeable about and sensitive to issues of interpersonal violence. Local resource registries may maintain a list of service providers (including health care practitioners) who specialize in working with survivors.

While some survivors may disclose past abuse as a lead-up to asking for a referral to specialized

Health care providers should not assume that all issues have been dealt with in one or two discussions; rather, they should check in with their clients throughout each appointment and make repeated invitations for feedback.
counselling or support services, it is a mistake to assume that all survivors who disclose need or want to be referred to a mental health practitioner. By offering a referral before exploring the survivors’ intentions, practitioners may give the impression that they think they know what is best for the individual or do not want to deal with the disclosure. An immediate referral to a mental health practitioner, regardless of whether the client is having difficulties related to past abuse, can feel like a clear statement that the clinician has judged the survivor to be “not okay.” Under many circumstances, raising the issue of referral to a mental health practitioner may best be postponed to a later interaction so that practitioners can reinforce their acceptance of the survivor after the disclosure.

Section 5.10 – Community resources for survivors and health practitioners

A preferable response to disclosure is for practitioners to ask about the presence and effectiveness of supports (e.g., friends, family, counsellor, spiritual advisor, or self-help group) available to the survivor. Such questioning gives the practitioner information about the survivor’s current resources and helps identify gaps. Questions such as “To whom do you turn for support?” or “Do you have enough support in your life?” can help assess the individual’s situation. Further questions can help the practitioner make survivors aware of the organizations in the community that offer information, support, and other services to survivors:

I needed to be reminded of resources and also that it was okay for me to call and use the resources. I needed permission to get the support I need. (Man survivor)

8.6 Responses to avoid following a disclosure

There are, unfortunately, instances when health care practitioners fail to respond sensitively to a disclosure. This failure often leaves the survivor who has risked sharing deeply personal information feeling more distressed.

TABLE 5
Components of an effective response to disclosure

<table>
<thead>
<tr>
<th>After hearing a disclosure of past abuse, the clinician should:</th>
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<tbody>
<tr>
<td>• Accept the information</td>
</tr>
<tr>
<td>• Express empathy and caring</td>
</tr>
<tr>
<td>• Clarify confidentiality</td>
</tr>
<tr>
<td>• Normalize the experience by acknowledging the prevalence of abuse</td>
</tr>
<tr>
<td>• Validate the disclosure</td>
</tr>
<tr>
<td>• Address time limitations</td>
</tr>
<tr>
<td>• Offer reassurance to counter feelings of vulnerability</td>
</tr>
<tr>
<td>• Collaborate with the survivor to develop an immediate plan for self care</td>
</tr>
<tr>
<td>• Recognize that action is not always required</td>
</tr>
<tr>
<td>• Ask whether it is a first disclosure</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>At the time of disclosure or soon after:</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Discuss the implications of the abuse history for future health care and interactions with clinician</td>
</tr>
<tr>
<td>• Inquire about social support around abuse issues</td>
</tr>
</tbody>
</table>
Negative responses (such as ignoring the disclosure, disbelief, denial of the negative impact of the abuse, or telling a survivor to “just get over it”) are both painful and silencing:

_He told me that I should just get over this and move on._ (Woman survivor)

_Don’t push the person and be really aware not to use the “shoulds,” like “You should call the crisis line.”_ (Woman survivor)

Men survivors also cautioned against minimizing the effects of female-perpetrated abuse. Viewing the survivor as lucky to have had such an early introduction to sex or perceiving the abuse as merely a sowing of wild oats was very damaging.

### 8.7 Legal and record-keeping issues

**Legal obligations.** In our studies, we use the term _disclosure_ to refer to survivors telling health care practitioners that they have a history of childhood abuse, as distinct from _task-specific disclosure_, which occurs when individuals identify discomfort or difficulty with all or part of a specific examination or treatment. With the exception of this section, when we speak about disclosure in this handbook, we are referring strictly to _adults_ revealing a history of _past abuse_.

Health care practitioners do not have a legal obligation to report _past_ child abuse disclosed by an _adult_, unless, in disclosing his or her own experience, an individual identifies a child who may be currently in need of protection (e.g., if a male patient who was abused by a family member tells the practitioner that he has reason to believe that the same perpetrator is continuing to abuse children).

In contrast, all Canadian jurisdictions, except Yukon, have laws that mandate a duty to report cases of suspected abuse or neglect of children to child welfare agencies or to police.103,127 Although the definitions of a child and the definitions of a child at risk vary somewhat among individual provinces,127 these same laws require that all cases of suspected abuse of children (under the age of majority) be investigated by the appropriate child welfare service to determine whether the children are in need of protection. When the suspicions are substantiated, child welfare authorities are mandated to intervene.

**Responses to avoid after a disclosure**

Survivors identified the following responses as clearly not helpful:

- Conveying pity (e.g., “Oh, you poor thing”).
- Offering simplistic advice (e.g., “Look on the bright side,” “Put it behind you,” “Get over it,” or “Don’t dwell on the past.”).
- Overstating or dwelling on the negative (“A thing like that can ruin your whole life”).
- Smiling (while you may hope that your smile conveys compassion, a neutral or concerned expression is more appropriate).
- Touching the person without permission even if you intend it as a soothing gesture.
- Interrupting (let the individual finish speaking).
- Minimizing or ignoring the individual’s experience of abuse, the potential impact of past abuse, or the decision to disclose (e.g., “How bad could it be?”; “I know a woman that this happened to and she became an Olympic gold medalist,” “Let’s just concentrate on your back pain,” or “What’s that got to do with your sprained ankle?”).
- Asking intrusive questions that are not pertinent to the examination, procedure, or treatment.
- Disclosing your own history of abuse.
- Giving the impression that you know everything there is to know on the subject.

If clinicians think that they have inadvertently responded to the disclosure in an inappropriate way, or if the patient’s nonverbal feedback suggests a negative reaction to their initial responses, they should immediately clarify the intended message and check with the survivor for further reaction.
A guiding principle underlying the Framework is that the collection, use, and disclosure of health information are to be done on a need-to-know basis and with the highest degree of anonymity possible under the circumstances. Furthermore, the Framework understands privacy as a consent-based right and, unless otherwise stated in legislation, an individual’s consent must be obtained for any collection, use, and disclosure of personal health information. As well, Alberta, Saskatchewan, Manitoba, and Ontario have enacted provincial legislation that addresses the collection, use, and disclosure of personal health information by health care providers and health care organizations.

Documenting a history of abuse. Some of the survivors in our studies specifically asked their health care providers not to document past abuse. In responding to this request, a practitioner needs to balance the patient’s right to privacy and legal reporting requirements. While not reporting suspected abuse of a child clearly contravenes the intent of the law, the same is not true of past abuse of a person who is currently an adult. Both practitioner and survivor participants concluded that it is important for clinicians to discuss with their patients how documentation of past abuse might be done while still protecting their privacy. Survivors, for example, might agree to a chart note that states they have a history of abuse but provides no further details. Privacy may, however, be an issue when working with patients whose care is being paid for by a third party (e.g., insurance companies, employee assistance programs, or workers’ compensation). Clinicians involved in fee-for-service practices are urged to pay particular attention to the reporting that is required of them.

Health care practitioners are further urged to consider the possible ramifications of sharing information about patients’ histories of abuse.

It is the responsibility of all health care practitioners to know the legal requirements for reporting child abuse and neglect in their jurisdiction.
when referring them to other health care practitioners:

On many occasions when I’ve been referred to a specialist, it has been noted in the referring letter/form that I have a history of abuse. Too often the referring [practitioner] assumes it is sexual abuse although I have never specified. I have learned the hard way that this information is not seen only by the [receiving health care practitioner] ... but is also read by some of the staff at the clinic I am going to. I don’t want my history of abuse broadcast to the world so I now ask the referring clinician to state that a sensitive approach to any physical examination is required rather than disclose my abuse history. If the referring doctor needs to know, I am in a position to disclose or not and to only that person. (Woman survivor)

Documentation about past abuse may have legal implications for clients who are (or may in the future be) involved in a court case. If, for example, a client chooses to press criminal charges or launches a civil action against an abuser, or if civil litigation follows a motor vehicle crash, relevant health records may be subpoenaed. Sometimes the records will be sought to support the client’s case; however, in other instances, they might be used to challenge the client’s credibility or account of events. Regarding consent for the provision of medical records to insurance companies, survivors should be advised that they have the option of sharing all or only specific portions of their record. Health care practitioners are strongly encouraged to seek legal advice in situations where a client’s health record is requested by a third party before taking any action.

Section 5.8 – Collaborative service delivery

8.8 Questions for reflection

- Does my environment foster a sense of safety for potential disclosure?
- Do my clients trust me enough to disclose? Are there any steps I could take to increase their feelings of trust and safety?
- How do I want to integrate routine inquiry about child sexual abuse? Do I have a “script” that feels natural to me?
- How would I feel if a client disclosed a history of child sexual abuse? Are my reactions different for males and females? How would I know whether my reactions are helpful for my clients?
9 Summary and Concluding Comments

9.1 Clinicians’ contributions to survivor’s healing from childhood sexual abuse

I think that we’re talking about really long-term partnerships with a number of medical people ... maybe a physiotherapist, a psychotherapist, a family doctor. We need these nuclei of support, and they need to be in touch with each other, and I have that, so I feel like I have a network of support. (Woman survivor) 143p.259

All forms of violence and abuse can leave an individual feeling disempowered and disconnected from others. Healing from abuse involves re-empowerment and reconnection with self and others.81 Because the harm of abuse occurs in the context of relationships and because it affects individuals’ ability to relate with others, healing can only occur in relationships. Relationships with caring others provide the substrate – the nutrient medium – for healing the parts of the self that were damaged by past trauma. The absence of trusting relationships leaves survivors isolated in their shame. Through engagement with others, survivors can learn to rebuild their basic capacities for autonomy, trust, and intimacy.81 Health care practitioners can be allies in that process by offering effective and sensitive health care in the context of genuine human connection. They can also facilitate reconnection by helping survivors learn about their bodies and how they function in health and illness.

9.2 Sensitive Practice and patient-centred care

Some would argue that the Sensitive Practice paradigm is redundant – that client-centred care, by definition, incorporates all that this Handbook describes. However, we have a different perspective. While the Handbook includes neither all possible ways that practitioners can be sensitive to survivors nor all of the ways that interpersonal violence can affect an individual’s health or health care experiences, we have come to see Sensitive Practice as a refinement or “fine tuning” of patient-centred care. If all practitioners were knowledgeable about the association between abuse and health, then Sensitive Practice might not be necessary. Unfortunately, many curricula devote little attention to violence and abuse and their implications for health and health care,169 and some practitioners remain convinced that our health care system cannot afford the few extra minutes it takes to enact Sensitive Practice. We suggest that, especially in light of the pressures on the health care system, failure to practice sensitively is tantamount to abdicating our ethical responsibility to do no harm.

Thus, we encourage all health care providers to become more aware of the effects of violence and abuse and to ensure that their words and behaviour communicate this understanding in a sensitive way. By fine-tuning our patient-centred approach, we will make greater strides in helping patients become healthier and better functioning members of society. Although a practitioner’s contributions to an individual’s healing are not always measurable, survivors have reminded us that the trust and safety that allow (re)connection within a strong therapeutic relationship can be hugely helpful to them. And the possibility that the survivor will be further empowered to make gains, however slow, holds bright promise:

So, what we have is a relationship of ... mutual give and take ... [The clinician] gives me a lot of responsibility; I give her a lot of information; we negotiate how best to work [together] to help me to fulfil my needs and to let me have power over my own life. (Woman survivor) 143p.260
This *Handbook* is informed by two multi-disciplinary, multisite research studies that employed grounded theory and action research methods. The overall intent of the project has been to facilitate a process by which childhood sexual abuse survivors and health care practitioners collaborate to develop practice knowledge that influences health care. In the first study, we explored women survivors’ experiences of physical therapy, consulted with survivors, physical therapists and physical therapy students to develop guidelines for Sensitive Practice, and summarized the results of the study in a handbook. In the second study, we addressed gaps and questions from the first project, by asking men and women survivors about their experiences with and ideas about Sensitive Practice for a wide range of health care practitioners (including but not limited to physicians, nurses, nurse practitioners, oral health practitioners, massage therapists, complementary therapy practitioners, and other health care practitioners without special training in mental health/psychiatry or psychotherapy). We then engaged health care practitioners and survivors in a dialogue about Sensitive Practice and the creation of this second edition of the *Handbook*.

The first research project was conducted in three phases. In the first phase, 27 adult female childhood sexual abuse survivors in Saskatchewan and Ontario were interviewed. The women were between 19 and 62 years of age and from a broad range of educational backgrounds, professions, and income levels; 26 identified themselves as Caucasian and one woman self-identified as Métis. All had been referred for physical therapy in either inpatient or outpatient clinics; four had declined to see physical therapists. Transcripts of the interviews were analyzed and the central themes identified.

In the second phase of the project, working groups of four survivors and four physical therapists in each group met in Saskatoon, Saskatchewan, and Waterloo, Ontario, four to six times over six months to refine the themes into principles and guidelines for Sensitive Practice. In the final phase of the project, the information from the interviews and working groups was used to draft the first edition of the *Handbook*. Approximately 200 survivor participants, other survivors, physical therapists, physical therapy students, and counsellors across Canada commented on successive drafts in writing or during focus groups. This lengthy and broad consultative process was intended to ensure the clinical relevance of the first edition of the *Handbook*.

In our second study, we focused on survivors’ experiences with all types of health care providers. We conducted individual interviews with 49 men survivors and talked with one group of nine men. Interviews were conducted in six provinces (British Columbia, Alberta, Saskatchewan, Ontario, Nova Scotia, and New Brunswick). We also conducted interviews with 19 women survivors in Saskatchewan and Ontario. Repeated efforts to recruit Aboriginal survivors and survivors of colour were not overwhelmingly successful. Eight men and one woman self-identified as Aboriginal (Métis or First Nations). Participants ranged in age from 24 to 62 years. Participants were from a broad range of educational backgrounds, professions, and income levels.

As in the first study, themes from these interviews served as the starting point for two working groups in each of two cities. In Winnipeg, one working group was made up of four male survivors and three nurses and nurse practitioners and the other working group consisted of four male survivors and four physicians. In Saskatoon, one working group included four female survivors and three nurses and the other was made up...
of three female survivors and four physicians. The groups met three or four times over four months. Interview findings and recommendations from the working groups in both studies were incorporated into a draft of the second edition of the Handbook that was sent to all participants. Feedback was incorporated into draft 2 and was sent to 110 health care practitioners representing a wide range of health disciplines and perspectives (including those from academic training programs, professional associations, regulatory/licensing bodies and various practice settings). Feedback was received from 56 consultants, including aroma therapists, chiropractors, dentists, dental hygienists, dental assistants, individuals teaching and researching in the area of kinesiology and sport, massage therapists, mental health practitioners, midwives, naturopathic doctors, nurses, nurse practitioners, physicians, physical therapists, occupational therapists, and reiki practitioners. Draft 3 was developed from this consultation and was used for further consultations with six focus groups of health care practitioners and students. In total, approximately 200 survivors and health care practitioners from across Canada participated in this consultation process.
Because childhood sexual abuse is often unreported in childhood or adolescence, adult retrospective studies are the most common source of prevalence estimates. The most current and reliable lifetime prevalence estimates are that as many as one third of women and 14% of men are survivors of childhood sexual abuse. Accurate accounting of the occurrence of childhood sexual abuse is hampered by methodological issues related to reporting barriers (e.g., shame, guilt, self-blame, fear, etc.), definitional controversies, population sampled (community vs. clinical), method of data collection (e.g., self-report questionnaire vs. interview), response rates, and the number of questions researchers ask about childhood sexual abuse. This helps to explain the wide range in reported results and suggests caution when interpreting results and making cross-study comparisons.

Large community-based studies of the incidence and prevalence of childhood sexual abuse among children and youth are rare, with the most comprehensive one being a telephone survey of 2,000 U.S. residents (aged 10-16 years) done by David Finkelhor and Jennifer Dziuba-Leatherman. These authors report that in the year preceding the interview, 3.2% of girls and 0.6% of boys had experienced contact childhood sexual abuse, which was defined as “a perpetrator touching the sexual parts of a child under or over the clothing, penetrating the child, or engaging in any oral-genital contact with the child.” In the overall sample, the combined prevalence of attempted and completed childhood sexual abuse categories was 10.5%.

Community-based probability samples typically find that 12%-35% of women and 4%-9% of men have had an unwanted sexual experience before the age of 18 years. After adjusting for sample-related variation, response rates, and differences in definitions across 16 cross-sectional community sample surveys, Kevin M. Gorey and Donald R. Leslie determined that the prevalence of childhood sexual abuse was 16.8% for women and 7.9% for men. In his review of large community-based studies in 19 countries around the globe, Finkelhor found that the prevalence of childhood sexual abuse was 7%-36% for females and 3%-29% for males, indicating that childhood sexual abuse is an international problem and has been found in every region where it has been studied. More recent studies in non-Western countries confirm this. David Murray Fergusson and Paul E. Mullen, after examining community-based prevalence from several countries, concluded that between 15% and 30% of females and between 3% and 15% of males report exposure to some form of unwanted sexual attention in childhood. This is consistent with John Briere and Diana Elliot’s recent study, which found 32.3% of women and 14.2% of men reported sexual abuse in childhood. In the latter work, 21% of adults who reported histories of childhood sexual abuse also experienced physical maltreatment.

The prevalence of childhood sexual abuse is even higher among individuals with disabilities. A review of literature regarding children with disabilities, defined as “the full spectrum of physical, mental, and emotional impairment,” cited research that reported children with disabilities are almost twice as likely to be neglected, 1.6 times more likely to be physically abused, and 2.2 times more likely to be sexually abused than are children without disabilities. Other studies report even higher rates of sexual abuse for children with disabilities.
Some of the common problems experienced by childhood sexual abuse survivors are summarized in Table 6 using David Finkelhor and Angela Browne’s conceptualization of the traumagenic dynamics of childhood sexual abuse. These dynamics describe the impact that abuse-related behaviours, events, and experiences can have on their victims.

### TABLE 6
Traumagenic dynamics of childhood sexual abuse

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Dynamics</th>
<th>Possible Manifestations</th>
</tr>
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| Traumatic Sexualization | - Rewarding a child for sexual behaviour may heighten the salience of sexual issues for him or her  
- Receiving attention and affection for sex can affect a survivor’s ability to achieve a healthy sense of love and belonging  
- Sexual parts of the child may be fetishized  
- Abuse may impart misconceptions about sexuality (e.g., sexual identity, sexual orientation, sexual behaviour, and sexual morality)  
- Sexual activities may become conditioned to negative emotions and memories, creating an aversion to sex or intimacy | - Avoidance of all things sexual  
- Preoccupation with sexual matters or compulsive sexual behaviours  
- Precocious sexual activity  
- Aggressive sexual behaviours  
- Promiscuity  
- Prostitution (making use of or working in the sex trade)  
- Sexual dysfunctions (e.g., lack of desire, difficulty with sexual arousal, inability to experience orgasm, and avoidance of sexual intimacy) |
| Betrayal                | - Childhood sexual abuse manipulates a child’s vulnerability, violates the expectation that others will provide care and protection, and may interfere with the ability to trust  
- The child’s autonomy and wellbeing are disregarded, which may affect the sense of self  
- Deep-seated guilt and shame develop because children believe something bad about them caused the abuse  
- Profound sense of grief over lost innocence or the “perfect” or “normal” family; depression  
- Extreme anxiety or fear, which engenders a sense of dependency | - Overdependence or clingingness  
- Vulnerability to subsequent abuse and exploitation  
- Failure to accurately judge the trustworthiness or motives of others, leading to subsequent abuse and exploitation and/or inability to protect one’s own children from abuse  
- Social withdrawal, isolation, and/or avoidance of intimate relationships  
- Chronic relationship difficulties  
- “Acting out” behaviours (e.g., aggression, delinquency, risk-taking, etc.) |
<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Dynamics</th>
<th>Possible Manifestations</th>
</tr>
</thead>
</table>
| Stigmatization  | • Abuser(s) and others blame or denigrate the victim engendering a sense of shame or guilt  
• The abuser and others pressure child for secrecy  
• The victim feels “damaged,” “abnormal,” “bad,” which may contribute to a distorted sense of self and lowered self-esteem |
|                 |                                                                          | • Dysphoria or chronic depression  
• Stigmatization, isolation, and marginalization may contribute to substance abuse  
• Criminal behaviour  
• Failure to care for oneself (e.g., risk-taking behaviours, poor hygiene, poor health practices)  
• Self-harm or self-mutilation |
| Powerlessness    | • Unwanted invasion of one’s body or personal space can interfere with the establishment and maintenance of healthy boundaries and increase risk of repeated victimization  
• Abuser(s) may use violence, threats, trickery, or bribery to involve their victim  
• If others do not believe and respond appropriately to disclosure of abuse, an individual may develop a lowered sense of efficacy  
• Some victims develop a high need for personal control and may even identify with the abuser |
|                 |                                                                          | • Hyper-arousal (i.e., chronic anxiety, phobias, tendency to startle easily, irritability, poor sleep)  
• Intrusion (e.g., flashbacks during waking states, traumatic nightmares during sleep)  
• Constriction (dissociation to endure danger that one is unable to fight off or escape) -alters perception, sensation, and time sense and may result in avoidance of reminders of the trauma, emotional numbing/blunting, detachment, and an inability to experience joy  
• Stress-related disease and illness; chronic and/or vague somatic problems |

Adapted from Finklehor and Browne\(^6\) with permission of D. Finkelhor.
Acute Stress Disorder (ASD)

Reprinted from The Diagnostic and Statistical Manual of Mental Disorders (4th ed., Text Revision) with permission of the American Psychiatric Association.

A. The person has been exposed to a traumatic event in which both of the following were present:

- the person experienced, witnessed, or was confronted with an event or events that involved actual or threatened death or serious injury, or a threat to the physical integrity of self or others
- the person’s response involved intense fear, helplessness, or horror

B. Either while experiencing or after experiencing the distressing event, the individual has three (or more) of the following dissociative symptoms:

- a subjective sense of numbing, detachment, or absence of emotional responsiveness
- a reduction in awareness of his or her surroundings (e.g., “being in a daze”)
- derealization
- depersonalization
- dissociative amnesia (i.e., inability to recall an important aspect of the trauma)

C. The traumatic event is persistently re-experienced in at least one of the following ways:

- recurrent images, thoughts, dreams, illusions, flashback episodes, or a sense of reliving the experience
- distress on exposure to reminders of the traumatic event

D. Marked avoidance of stimuli that arouse recollections of the trauma (e.g., thoughts, feelings, conversations, activities, places, people).

E. Marked symptoms of anxiety or increased arousal (e.g., difficulty sleeping, irritability, poor concentration, hypervigilance, exaggerated startle response, motor restlessness).

F. The disturbance causes clinically significant distress or impairment in social, occupational, or other important areas of functioning or impairs the individual’s ability to pursue some necessary task, such as obtaining necessary assistance or mobilizing personal resources by telling family members about the traumatic experience.

G. The disturbance lasts for a minimum of 2 days and a maximum of 4 weeks and occurs within 4 weeks of the traumatic event.

H. The disturbance is not due to the direct physiological effects of a substance (e.g., a drug of abuse, a medication) or a general medical condition, is not better accounted for by Brief Psychotic Disorder, and is not merely an exacerbation of a pre-existing Axis I or Axis II disorder.

Posttraumatic Stress Disorder (PTSD)

Reprinted from The Diagnostic and Statistical Manual of Mental Disorders (4th ed., Text Revision) with permission of the American Psychiatric Association.

A. The person has been exposed to a traumatic event in which both of the following were present:

1. the person experienced, witnessed, or was confronted with an event or events that involved actual or threatened death or serious injury, or a threat to the physical integrity of self or others.
(2) the person’s response involved intense fear, helplessness, or horror. Note: In children, this may be expressed instead by disorganized or agitated behaviour.

B. The traumatic event is persistently reexperienced in one (or more) of the following ways:

(1) recurrent and intrusive distressing recollections of the event, including images, thoughts, or perceptions. Note: In young children, repetitive play may occur in which themes or aspects of the trauma are expressed.

(2) recurrent distressing dreams of the event. Note: In children, there may be frightening dreams without recognizable content.

(3) acting or feeling as if the traumatic event were recurring (includes a sense of reliving the experience, illusions, hallucinations, and dissociative flashback episodes, including those that occur on awakening or when intoxicated). Note: In young children, trauma-specific reenactment may occur.

(4) intense psychological distress at exposure to internal or external cues that symbolize or resemble an aspect of the traumatic event.

(5) physiological reactivity on exposure to internal or external cues that symbolize or resemble an aspect of the traumatic event.

C. Persistent avoidance of stimuli associated with the trauma and numbing of general responsiveness (not present before the trauma), as indicated by three (or more) of the following:

(1) efforts to avoid thoughts, feelings, or conversations associated with the trauma

(2) efforts to avoid activities, places, or people that arouse recollections of the trauma

(3) inability to recall an important aspect of the trauma

(4) markedly diminished interest or participation in significant activities

(5) feeling of detachment or estrangement from others

(6) restricted range of affect (e.g., unable to have loving feelings)

(7) sense of a foreshortened future (e.g., does not expect to have a career, marriage, children, or a normal life span)

D. Persistent symptoms of increased arousal (not present before the trauma), as indicated by two (or more) of the following:

(1) difficulty falling or staying asleep

(2) irritability or outbursts of anger

(3) difficulty concentrating

(4) hypervigilance

(5) exaggerated startle response

E. Duration of the disturbance (symptoms in Criteria B, C, and D) is more than 1 month.

F. The disturbance causes clinically significant distress or impairment in social, occupational, or other important areas of functioning.

Specify if: Acute: if duration of symptoms is less than 3 months
Chronic: if duration of symptoms is 3 months or more

Specify if: With Delayed Onset: if onset of symptoms is at least 6 months after the stressor

**Disorders of Extreme Stress not Otherwise Specified (DESNS)**

Judith Herman has challenged the ability of the diagnosis of posttraumatic stress disorder (PTSD) to capture the full range of human response to trauma. She and others (e.g.,) suggest that it is more accurate to think about human responses to trauma as a spectrum anchored at one end by an acute stress reaction that resolves on its own without treatment, and on the other by what Herman calls “complex posttraumatic stress disorder,” with “classic or simple” PTSD residing somewhere between the two.

When the **DSM IV** was being developed, a field trial was completed to explore whether the construct of complex PTSD, also termed **disorders of extreme stress, not otherwise specified** (DESNS) should be included as a diagnosis separate from
PTSD. Although many argue that the field trial and other more recent studies support the legitimacy of the DESNOS diagnosis, DESNOS was not included in the DSM IV. However, the categories of symptoms included in the conceptualization of DESNOS were listed under the “Associated and Descriptive Features of PTSD.”

In the most recent version of DSM-IV-TR, the authors state that the following constellation of symptoms may be associated with PTSD, and are generally seen when the stressor involves interpersonal trauma such as childhood sexual or physical abuse or domestic battering:

- impaired affect modulation;
- self-destructive and impulsive behaviour;
- dissociative symptoms;
- somatic complaints;
- feelings of ineffectiveness;
- shame, despair, or hopelessness;
- feeling permanently damaged;
- a loss of previously sustained beliefs;
- hostility;
- social withdrawal;
- feeling constantly threatened;
- impaired relationships with others;
- a change from the individual's previous personality characteristics.11p.465
APPENDIX E: Sample Introduction to a Facility

Survivors pointed out that they are unfamiliar with the scope of practice for many health care practitioners and much of what is involved in examination and treatment. The following is an information sheet for new patients developed by survivors and clinicians to provide an understandable introduction to an out-patient physical therapy facility. Health care providers are encouraged to work with patients to develop introductions to their practice and facilities similar to this physical therapy example.

Suggestions for clients at out-patient physical therapy facilities

Welcome to physical therapy! We are glad to work with you. Physical therapy will include an assessment and treatment by the physical therapist. Direct and open communication between the client and the therapist is important. Below is a list of suggestions that may help you at physical therapy.

You have the right to choose a male or female physical therapist.
- If you know this is important for you, please tell us when you book your first appointment.
- If you decide later in treatment that you would rather work with a therapist of a different gender, you may tell us then too.
- If we are unable to book you with your choice of a male or female therapist, we may refer you to a facility that can.

You can choose to have someone accompany you during your physical therapy appointments. This person can be:
- a family member or friend.
- a staff member from the clinic.

Physical therapy works best when you and your therapist work as a team. For example, your physical therapist will explain your treatment to you. Please tell your physical therapist if:
- you are not comfortable with the treatment.
- you do not understand the treatment or language your therapist is using.
- you do not agree with the treatment.

Also, physical therapy works best when you talk to your physical therapist about how the treatment is working (or not working) for you. The more you are able to tell your physical therapist, the better he or she will be able to help you.

We will do our best to ensure your privacy.
- Your physical therapist may need you to wear a gown for some treatments. If you would prefer to bring loose fitting clothing from home, please tell your physical therapist.
- In some cases it is necessary to change your clothing for your treatment: you will have privacy to change your clothing.
- Please tell us if you would like the curtains drawn around your treatment table during any part of treatment.

Physical therapy involves touch and movement of your body. Tell your physical therapist if:
- certain parts of your body are sensitive to touch or movement.
- you are nervous about touch.
- there is something your physical therapist can do to make you more comfortable.

You have the right to stop treatment at any time, during or after a session. Reasons people might stop treatment may include:
- discomfort during treatment.
- deciding to try a different type of medical care.

If you decide to try a different type of care, your physical therapist may be able to give you the name of someone she or he thinks can help you.

Above all, we want you to notice an improvement in your health.
Consent Form for Examination and Treatment by a Physical Therapist

I am about to be examined and treated by a physical therapist and her/his assistants.

In order for me to be properly examined or treated, I will need to wear shorts and a T-shirt. The physical therapist will need to observe my body while it is still and while it is moving. It will be necessary for the therapist to touch and move my body in assessment and treatment. Should I feel uncomfortable about the assessment and treatment process at any time, I can inform the physical therapist and request that assessment and/or treatment be stopped. I can have someone else in the room with me; either a friend or relative, or someone else from the clinic, if available. In the latter case, I can choose the gender of this person.

I will need to tell the physical therapist about my health problems, both past and present. The therapist will ask my permission to contact my doctor if he or she finds any new problems. I am aware that all information I disclose and all information that will be charted is confidential.

Physical therapy treatment may involve: ______________(plain language, be specific). Soreness after treatment is common because joints and muscles are stretched. If I have any other symptoms, I will tell my physical therapist.

My signature below indicates that I understand all of the above information.
A brief history of Aboriginal Peoples

Abundant literature describes the history of the First Nations, Métis and Inuit in Canada. An abbreviated list of recommended readings is found in the bibliography. The purpose here is to provide an overview as a starting point for health care practitioners working with First Nations, Métis, and Inuit peoples in the area of Sensitive Practice.

Aboriginal Peoples constitute a diverse population in Canada. There are several terms that have been introduced in the literature in an attempt to categorize these populations into one group, such as aboriginal, native, Indian, and indigenous. The Constitution of Canada uses the term Aboriginal to include Status and Non-Status Indians, Métis and Inuit. Status Indians are those whose ancestors signed treaties; Non-Status Indians are those whose ancestors refused to sign treaties or were absent at the time of the signing. A subpopulation of Non-Status Indians was also created through loss of treaty rights for various reasons such as serving in the armed forces, voting, obtaining a postsecondary degree, and, for Status Indian women, marrying non-Aboriginal men. For the most part, members of this subpopulation have regained their treaty rights through a revision in the Indian Act in 1984 (Bill C-31).

Our knowledge of the history of indigenous people in Canada prior to the arrival of the Europeans is very limited. The majority of information has been gleaned through the sciences of archaeology and anthropology. The most commonly held theory in the Western world is that the ancestors of the First Nations came from Asia over the Bering Strait. *Time immemorial*, a phrase often used by First Nations to describe how long they have been here, has been roughly translated to mean between 50,000 and 15,000 BC. There have been three separate times when the Bering Strait could have been used as a land bridge, and there are theories arguing that there were three distinct migrations. On the basis of archaeological findings, it appears that, through multiple generations, the first wave of people travelled down the Pacific coast into South America. As the glaciers retreated, some headed back up north. The second wave, the Athapascans (Dene), stayed in the north, but began to move south following a volcanic eruption. The third wave, the Inuit, spread eastward through the north.

There have been several attempts to categorize the First Nations people of Canada. Linguistics is one common method, and there are 11 different language families: Algonquian, Athapaskan, Eskimo-Aleut, Haida, Tlingit, Siouan, Tsimshian, Wakashan, Salishan, Kutenai, and Iroquoian. It has been hypothesized that around the time of European contact there were between 50 and 60 languages, but the most commonly spoken languages today are Cree, Ojibway, and Inuktitut. Many First Nations are diligently working to save their languages. Another method of categorization is culture areas, and these areas are based on geography and a group of people sharing similar cultures. These culture areas are: Arctic, Western Subarctic, Eastern Subarctic, Northeastern Woodlands, Plains, Plateau, and Northwest Coast. It is interesting that these geographical culture areas closely resemble the geographical separation according to linguistics. Today there are more than 610 First Nations communities in Canada, and the total population, living both on and off reserve, is more than 733,000.

The time following contact with Europeans brought many changes to First Nations peoples in Canada, including the creation of an entirely different people: the Métis. The Métis were primarily the offspring of First Nations women and French men. The term *half-breed* was more
often used to describe children of First Nations women and Scottish or English men. The Métis of today define themselves according to the following definition adopted by the Métis National Council, the national governmental organization representing the Métis: “Métis means a person who self identifies as Métis, is of Historic Métis Nation ancestry, is distinct from other Aboriginal peoples and is accepted by the Métis Nation.” The “Historic Métis Nation” means the Aboriginal people then known as Métis or Half-Breeds who resided in the Historic Métis Nation Homeland, the area of land in west central North America used and occupied as the traditional territory of the Métis or Half-Breeds. The Métis National Council estimates that there are between 350,000 and 400,000 Métis in Canada.

The Inuit are peoples who live in the Arctic regions of Canada, Alaska, and Greenland. They have very similar cultural and physical characteristics despite the wide geographical area in which they live. The Inuit have survived in one of the world’s harshest environments for more than 5,000 years. The areas are mostly coastal, consisting of shallow basins with rivers flowing through and many islands covered with permanent ice and mountain glaciers. The treeless shores provide no wind protection, temperatures are below freezing for eight or nine months of the year, and total precipitation is so slight that the area nearly qualifies as desert. According to the 2001 census, there are more than 45,000 Inuit in Canada, representing about 5% of the Aboriginal population. They are represented nationally by the Inuit Tapiriit Kanatami.

The residential school legacy

Between 1892 and 1969, approximately 135 residential schools were established to meet the treaty right to education. Although First Nations leaders wanted schools built on the reserves, the federal government decided that residential schools would be cheaper and entered agreements with the Roman Catholic Church, the Church of England, the Methodist Church, and the Presbyterian Church to operate the schools. The vast majority of these schools were in the western provinces and it is estimated that more than 150,000 students attended them. The Assembly of First Nations estimates that more than 105,000 survivors of residential schools are still alive today.

The premise of the residential schools was assimilation through education, religious indoctrination, and cultural degradation (teaching the children to be ashamed of their heritage). Physical, emotional, and sexual abuses were rampant and living conditions were often substandard. Former residents say that they were often hungry and that their parents brought them food on their weekend visits; others report being forced to steal food from the kitchens. The education the children received was also substandard. As late as the 1950s, more than 40% of the teaching staff at the schools had no professional training. Cultural degradation practices included physical and emotional abuse for speaking a traditional language, cutting students’ hair (hair has strong cultural and spiritual implications), imposing foreign religious practices, and intentionally separating students from visiting parents.

The residential school experiences continue to have a detrimental impact on Aboriginal communities today. These “intergenerational impacts refer to the effects of physical and sexual abuse that were passed on to the children, grandchildren, and great-grandchildren of Aboriginal people who attended the residential school system.” Some of these effects include:

- Alcohol and drug abuse;
- Past and ongoing physical, emotional, and sexual abuse;
- Low self-esteem;
- Dysfunctional families and interpersonal relationships;
- Parenting issues;
- Suicide;
- Teen pregnancy.

The Aboriginal Healing Foundation (AHF) was established in 1998 in response to the Royal
Health care systems

Health care is not a provision specifically addressed in the treaties between Canada and First Nations and Inuit. The only direct mention of health care can be found in Treaty 6 which was signed in the mid-prairies of Saskatchewan and Alberta in 1876 and reads, “That a medicine chest shall be kept at the house of each Indian Agent for the use and benefit of the Indians at the direction of such agent.” Subsequent court proceedings involving Treaty 6 have ruled that at the time the treaties were signed, the Chiefs were looking for the best possible agreement for their members, and within that understanding, the clause could mean the provision of any and all services necessary for continued health for First Nations. The intent of the medicine chest clause has been applied to all First Nations and Inuit peoples.

The federal government provides comprehensive health care services to First Nations and the Inuit through the First Nations and Inuit Health Branch (FNIHB). FNIHB provides direct care to on-reserve populations and reimburses the provincial and other health care agencies for services provided to off-reserve populations. There has been a recent shift in responsibility as First Nations are reclaiming some aspects of self-government. Health transfer payments to individual First Nations or Tribal Councils has allowed First Nations to administer the funding and given them the freedom to determine their own health needs and plan their programs accordingly. Non-Status and Métis people are left out of these arrangements and receive their health care within the provincial or territorial health system.

Aboriginal health beliefs

The most common health model found in the literature and the oral tradition of Aboriginal peoples is the medicine wheel model (see Figure 2). Actual medicine wheels are circular stone formations found in all parts of North America. The term medicine wheel has been borrowed from these stone structures and applied to the theory of health and other areas of Aboriginal traditions. The medicine wheel is a circle, which means there

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Recommended Readings and Resources – Aboriginal Peoples: Readings (especially the policy statements by Dr. Janet Smylie)
is no end and no beginning. The same could be said for one’s health status. The four areas of the wheel are intellectual, emotional, spiritual, and physical. Some Aboriginal people believe that all four areas have to be in balance if one is to be in an optimum state of health; in other words, if any of the four areas are out of balance, then the individual becomes ill. All four areas are also connected and interrelated, so that there is no distinction such as the separation between mind and body that is often found in Western health paradigms. Specific programs have been developed in many Aboriginal communities and organizations using the medicine wheel as the framework. An Internet search of the term medicine wheel reveals the diverse situations and disease entities to which this framework has been applied.

FIGURE 2
Medicine Wheel

Physical

Spiritual

Intellectual

Emotional
Dissociative Identity Disorder (DID) – which was previously called Multiple Personality Disorder – is a psychiatric condition which requires specialized training to diagnose correctly. The *DSM-IV-TR* criteria for DID include “the presence of two or more distinct identities or personality states (each with its own relatively enduring pattern of perceiving, relating to, and thinking about the environment and self).”

Researchers believe that DID is almost always associated with a history of severe child abuse and requires very specialized treatment by a multidisciplinary team. If a health care practitioner encounters a person who has been diagnosed with DID and is not already receiving treatment from a mental health team, a referral to and collaboration with such a specialized service is essential.
Appendices A to I

APPENDIX I: The Evidence Debate Pertaining to Inquiry about Interpersonal Violence

While the empirical evidence is clear about the high prevalence of childhood sexual abuse and links between childhood adversity and adult health problems, it is less clear about whether health care practitioners should routinely assess for current and past abuse/violence (usually including intimate partner (or domestic) violence and childhood abuse). Three recent systematic reviews from the United States, Britain, and Canada concluded that there is insufficient evidence to recommend routine screening for family and/or intimate partner violence. In contrast, the Intimate Partner Violence Working Group of the Society of Obstetricians and Gynaecologists of Canada (SOGC) and the Registered Nurses of Ontario both issued clinical practice guidelines endorsing routine assessment for intimate partner violence/woman abuse as standard practice. Furthermore, existing practice guidelines and recommendations for management of a number of conditions also call for an assessment of abuse history (e.g., 6,7, 8,12).

Section 2.5 – Childhood sexual abuse and health

This debate is pertinent to any discussion of making inquiries about a history of child sexual abuse. Exposure to childhood violence or abuse increases an individual’s risk for intimate partner violence in adulthood (e.g., 17). Because the two co-occur with some frequency, childhood abuse/violence and intimate partner violence may not be categorically discrete entities. This means that if an individual discloses intimate partner violence, there is also the possibility of past childhood abuse/violence.

Considerable expert opinion (including 57,153,181) disagrees with the findings of the systematic reviews cited above. Much of this disagreement centres around the distinction between inquiring about violence and screening (which by definition must meet strict requirements related to lack of symptoms, specificity, sensitivity, positive predictive value, negative predictive value, etc.). Those who oppose universal screening point to the “absence of any high quality evidence of the benefit and a similar lack of evidence that screening does not harm.”181p.163 Those who support routinely inquiring about violence point out that such inquiry does not equate to screening but rather, represents “asking questions about domestic violence during a health care contact.”181p.163 The SOGC reiterates this in its consensus statement on intimate partner violence screening:

Asking women about violence is not a screening intervention [emphasis added]: victims are not asymptomatic; disclosure is not a test result, it is a voluntary act, and the presence or absence of violence is not under the victims’ control; and most interventions required to protect and support survivors are societal, not medical.153p.366

Because the three systematic reviews looked only at those studies that met the criteria for “screening,” they considered only a small portion of the existing intimate partner violence research. For example, of the 806 abstracts that related to screening for intimate partner violence, only 14 met the inclusion criteria that Nelson and colleagues used; similarly, only two of the 667 abstracts on intimate partner violence intervention studies were considered.57 This led the Family Violence Prevention Fund’s Research Committee to conclude:

As a consequence of this overly narrow approach to what the most relevant research questions are, an important body of studies related to IPV [intimate partner violence] was not considered. The outcomes most closely focused on are harm, death, and disability. In contrast, most researchers in the field would expect that measurable benefits (desirable
We believe that the research on which this Handbook is based further supports the argument that routinely inquiring about a history of past abuse is not harmful to individuals and, if done in a sensitive and informed manner, is likely to lead to improved health for all patients.
Works Cited


Handbook on Sensitive Practice for Health Care Practitioners


Recommended Reading and Resources

Childhood sexual abuse and trauma: Readings


Pain: Readings


Childhood sexual abuse and trauma: Websites

National Clearinghouse on Family Violence has many resources available free of charge. Toll free telephone 1-800 267-1291; (613) 957-2938; Fax (613) 941-8930; http://www.phac-aspc.gc.ca/nc-cn

Canadian Association of Sexual Assault Centres is a pan Canadian organization of sexual assault centres in Canada. Their website has contact information and links for sexual assault centres across Canada. http://www.casac.ca/english/home.htm
Tamara’s House, a residential treatment centre for women survivors of childhood sexual abuse (Saskatoon, Saskatchewan) http://www.tamarashouse.sk.ca/


Men’s Resource Centre (Winnipeg, Manitoba) http://www.elizabethhill.ca/mrc.html

The Men’s Project (Ottawa/Cornwall Ontario) http://themensproject.ca

**Ritual abuse: Readings**


**Sexual abuse in sport: Readings**


**Critical cultural perspective: Readings**


Gustafson D.L. (Accepted). Beyond sensitivity and tolerance: Theoretical approaches to caring for newcomer women with mental health problems. In S. Guruge & E Collins (Eds.). *Working with women and girls in the context of migration and settlement*. Toronto, ON: Centre for Addiction & Mental Health.


**Aboriginal Peoples: Readings**


Aboriginal Peoples: Websites

Assembly of First Nations: http://www.afn.ca/
First Nations & Inuit Health Branch website: http://www.hc-sc.gc.ca/fnih-spni/index_e.html
Métis National Council: http://www.metisnation.ca/
Aboriginal Healing Foundation: http://www.ahf.ca/

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The goal of Sensitive Practice is to foster a sense of safety for patients. Although the principles and guidelines articulated in this *Handbook* and outlined in Tables 7 and 8 are based on studies with Canadian men and women with histories of childhood sexual abuse, they represent a basic approach to care that should be extended to all patients. By adopting the principles of Sensitive Practice as the standard of care, health care providers convey respect, support clients’ autonomy and right to participate in healthcare, and decrease the likelihood of inadvertently retraumatizing the survivors of abuse with whom they work knowingly or unknowingly.

**TABLE 7**

*Summary of principles of Sensitive Practice*

<table>
<thead>
<tr>
<th>Principle</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Respect</td>
<td>Acknowledging the inherent value of clients as individuals with unique beliefs, values, needs, and histories means upholding and defending their basic human rights and suspending judgment of them.</td>
</tr>
<tr>
<td>Taking time</td>
<td>Taking adequate time with patients ensures that they do not feel depersonalized or objectified.</td>
</tr>
<tr>
<td>Rapport</td>
<td>Developing and maintaining an interpersonal style that is professional, yet conveys genuine caring, promotes trust and a sense of safety.</td>
</tr>
<tr>
<td>Sharing information</td>
<td>Informing patients of what to expect on an ongoing basis and inviting them to ask questions and offer information and feedback helps reduce anxiety and promotes active engagement in their health care.</td>
</tr>
<tr>
<td>Sharing control</td>
<td>Seeking consent and offering choices enables the clinician to work <em>with</em> rather than <em>on</em> patients, and ensures that patients become full active participants in their own health care.</td>
</tr>
<tr>
<td>Respecting boundaries</td>
<td>Paying ongoing attention to boundaries and addressing difficulties that arise reinforces patients’ right to personal autonomy.</td>
</tr>
<tr>
<td>Fostering mutual learning</td>
<td>Fostering an environment in which information sharing is a two-way process encourages survivors to learn about their health and how to become active participants in their own health care. It also assists clinicians to learn how best to work with individuals who have experienced interpersonal violence.</td>
</tr>
<tr>
<td>Understanding nonlinear healing</td>
<td>Checking in with patients throughout each encounter and over time, and being willing to adjust their actions accordingly, enables caregivers to meet the needs of individuals whose ability to tolerate health care examinations and procedures vary over time.</td>
</tr>
<tr>
<td>Demonstrating awareness and knowledge</td>
<td>Showing that they are aware of interpersonal violence helps professionals foster a sense of trustworthiness and promotes an atmosphere in which patients are willing to work alongside their health care providers.</td>
</tr>
</tbody>
</table>
### TABLE 8
Summary of guidelines of Sensitive Practice

<table>
<thead>
<tr>
<th>Context of encounters</th>
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</table>
| **Administrative staff and assistants** | - Train all personnel about Sensitive Practice  
- Work with staff and assistants to establish a few “routine responses” that are survivor-friendly  |
| **Waiting areas** | - Keep patient informed of length of wait or invite patient to check intermittently  
- Provide printed materials related to interpersonal trauma  
- Provide and clearly identify washrooms  |
| **Privacy** | - Knock and wait for acknowledgement before entering  
- Have at least one soundproof examination or interview room  
- Problem-solve with patients to meet their needs for privacy and safety  |
| **Preparation of clients** | - Provide introductory information in plain language, both written and verbal  
- Negotiate with patient to identify needs and workable solutions  
- Encourage presence of support person or chaperone; agree upon roles for all parties  |
| **Encounters with patients** |  |
| **Introductions** | - Discuss and negotiate roles for patient and clinician prior to all examinations or treatments  
- Allow enough time to help individuals understand fully what you are doing  
- Do not assume the patient knows what is involved in an exam, treatment, or procedure  
- Seek consent in an ongoing way throughout the encounter  |
| **Clothing** | - Meet patient fully clothed before and after  
- Explain why removal of clothing is necessary  
- Discuss clothing requirements with patients and collaborate with them to find an agreeable solution  
- Minimize amount of clothing being removed and length of time patient must be disrobed  
- Provide gowns in a wide variety of sizes for all body types  
- Leave the room while the patient is changing  |
| **Task-specific inquiry** | - Inquire about patient’s past experiences, preferences, difficulties with the exam/ procedure  
- Inquire about how to increase the person’s comfort  
- Inquire about whether the patient thinks there is anything else that the clinician should know about  
- Repeat inquiry intermittently over time, and if body language suggests discomfort  |
| **General approach** | - Use task-specific inquiry to identify difficulties; problem-solve together to increase comfort  
- Monitor body language and follow up on signs of distress  
- Explain why positions for patient and clinician are necessary  |
| **Touch** | - Describe what is involved before and during the exam or treatment  
- Seek consent before beginning and when shifting from one part of the body to another  
- Encourage individuals to ask you to pause, slow down, or stop the examination or treatment at any time to lessen their discomfort or anxiety  
- When a presenting problem necessitates examination of areas of the body other than the site of the symptoms, explain the rationale  |
| **Genital, rectal exams and procedures** | - Acknowledge discomfort  
- Offer a running commentary about what you are doing  
- Minimize time the patient must remain in a subordinate position  
- Drape parts of the body not being examined  |
Oral and facial health care

- Agree on hand signals so that the patient can give instant feedback when verbal communication is not possible
- Problem-solve with the patient around difficulties with smell/feel of gloves, dental dam, body position, other task-specific difficulties
- Ensure that your comments to the patients about their oral health and behaviour during the appointment are offered in a nonjudgmental way
- Keep the length of appointments as short as possible; consider doing longer procedures over two or more appointments

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<th>Challenges in encounters</th>
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| Pain & disconnection from body | - Assess pain in systematic, nonjudgmental manner  
- Work with client to set realistic goals and determine appropriate referrals  
- Repeatedly invite individuals to focus on their bodies  
- Provide clear verbal and written instructions that the patient understands  
- Suggest a range of strategies to increase self-awareness |
| Non-adherence to treatment | - Explore all types of barriers with the patient and problem solve to identify workable solutions  
- Adapt treatment to fit patient  
- Create a “same-day” appointment for patients who frequently cancel appointments whenever possible |
| SAVE the situation | - Stop what you are doing and focus fully on the present situation  
- Appreciate and understand the person’s situation  
- Validate the person’s experience  
- Explore the next steps with the patient |
| Triggers and dissociation | - Examine list of common triggers and consider what can be avoided/accommodated  
- Become familiar with signs of a ‘fight or flight’ response  
- Work with patients who have been triggered to ground and reorient them  
- Normalize the experience  
- Ensure adequate follow-up |
| Anger and agitation | - Pay attention to personal safety  
- Adopt non-threatening body language  
- Negotiate and assure patients of your interest and concern |
| Disclosure |  |
| Responding effectively | Upon hearing a disclosure of past abuse:  
- Accept the information  
- Express empathy and caring  
- Clarify confidentiality  
- Normalize the experience by acknowledging the prevalence of abuse  
- Validate the disclosure and offer reassurance to counter feelings of vulnerability  
- Address time limitations  
- Collaborate with the survivor to develop an immediate plan for self care  
- Recognize that action is not always required  
- Ask whether it is a first disclosure  
| At the time of disclosure or in a future interaction:  
- Discuss the implications of the abuse history for future health care and interactions with clinician  
- Inquire about social support around abuse issues |