Introduction

In Western countries, prostate cancer (PCa) is the most common male cancer (Australian Government, 2013; Canadian Cancer Society, 2013; Centers for Disease Control and Prevention, 2013). Affecting 1 in 7 men, the incidence of PCa is increasing with an aging male population and improved screening (Canadian Cancer Society, 2013). Men are also living longer with PCa, and the death rate continues to be low amid increases in the number of men being diagnosed (Australian Government, 2013; Canadian Cancer Society, 2011; Centers for Disease Control and Prevention, 2013). In the absence of a known modifiable cause, for many men, PCa will be a chronic illness (Canadian Cancer Society, 2013; Jemal et al., 2009; Remzi, Waldert, & Djavan, 2004). Diverse morbidities can accompany PCa and its treatments, including erectile dysfunction and urinary incontinence; psychosocial supports are thus integral to the well-being of men who have PCa and their families (Arrington, 2010; Chapple & Ziebland, 2002; Charmaz, 1995; Fergus, Gray, & Fitch, 2002a, 2002b; Gannon, Guerro-Blanco, Patel, & Abel, 2010; Meuleman & Mulders, 2003; Navon & Morag, 2003a, 2003b; Oliffe, 2005, 2006; Wall & Kristjanson, 2005). Among an array of psychosocial oncology options, prostate cancer support groups (PCSGs) have prevailed as important community-based resources in Canada and other Western countries (Arrington, 2010; Arrington, Grant, & Vanderford, 2010; Gray et al., 1999; Gray, Fitch, Davis, & Phillips, 1996; Gray, Fitch, Davis, & Phillips, 1997; Gray, Fitch, Phillips, Labrecque, & Fergus, 2000b; Matsunaga & Gotay, 2005; Steginga et al., 2001; Zanchetta, Cognet, Xenocostas, Aoki, & Talbot, 2007). However, the sustainability of
volunteer-run PCSGs have been challenged by recruitment and leadership succession issues (Oliffe et al., 2008). Relatedly, PCSGs diverse connections to cancer agencies and health care providers can influence group viability. For example, health care providers who share information at PCSGs and/or refer patients to group meetings can advance the effectiveness and reach of PCSGs. While PCA specialists are an important health care provider subgroup, well positioned to link with PCSGs in a variety of ways the perceptions and practices of PCA specialists regarding PCSGs are poorly understood. In addressing the research question, what are Canada-based PCA specialists’ views about PCSGs? This study and article offers some empirical insights to address this knowledge gap and thoughtfully consider avenues for maximizing the potential of PCSGs.

Background

In contrast to the diversity in clinical psychosocial PCA programs, PCSGs are somewhat uniform in their provision of support, with both face-to-face and virtual groups focusing on (1) educational presentations from medical experts and/or (2) group discussions about treatment options, disease management, and/or health promotion (Oliffe, Gerbrandt, Bottorff, & Hislop, 2010; Seale, Ziebland, & Charteris-Black, 2006; Thaxton, Emshoff, & Guessous, 2005). Research indicates that men attend PCSGs to learn about their disease and treatment options and side effect management from other men who have direct experience, as well as from an array of health care providers (Arrington et al., 2010; Gray et al., 1997; Klemm, Hurst, Dearholt, & Trone, 1999; Manne, 2002; Oliffe et al., 2010; Seale et al., 2006; Smith, Crane, Byers, & Nelson-Marten, 2002; Thaxton, Emshoff, & Guessous, 2005). Women partners also attend PCSGs (Bottorff et al., 2008; Thaxton et al., 2005) and the group leaders can comprise health care providers and survivors. Compared with information provision, some have argued that PCSGs are less equipped to support men’s emotional distress (Arrington et al., 2010; Coriel & Behal, 1999; Weber & Sherwill-Navarro, 2005). Although experiencing significant psychosocial concerns in relation to being diagnosed, it has also been reported that men typically adopt an “instrumental” coping style oriented to action and focus on “functionality” more than “feelings” in the context of in-person and online PCSGs (Klemm et al., 2003; Oliffe et al., 2010; Seale et al., 2006). So while many men do share ordinarily private illness experiences at PCSGs (Oliffe et al., 2010), research indicates that emotional supports are secondary motivations for attendance (Thaxton et al., 2005; Voerman et al., 2007).

The vast majority of PCSG research has focussed on the benefits and barriers to attending group meetings. Previous work detailing benefits includes an interview study of 12 men—most of whom had leadership roles in PCSGs located at Winnipeg, Toronto, and Ottawa—reporting that men derived a sense of meaning and purpose through attending PCSGs (Gray et al., 1997). Similarly, a survey of men who attended a professionally led, Montreal-based PCSG reported that the sharing of PCA experiences gave men reassurance, helped alleviate anxiety, and provided a positive outlook and perception of being involved in their treatment (Gregoire, Kalogeropoulos, & Corcos, 1997). The benefits described in these studies are confirmed elsewhere wherein PCSGs have been reported as useful in mitigating the psychosocial aspects of cancer by conveying information, empowering men with PCa, enhancing and facilitating psychosocial adjustment, and helping men and their partners cope with life after a diagnosis and treatment of cancer (Arrington, 2003; Arrington et al., 2010; Cordova et al., 2003; Gray et al., 1997; Katz et al., 2002; Manne, 2002; Steginga et al., 2001; Voerman et al., 2007; Walker, 2005; Weber et al., 2004; Zhang et al., 2008).

In terms of barriers to attending PCSGs, an interview study with 34 Canadian men and their partners led Gray, Fitch, Phillips, Labrecque, and Fergus (2000a) to suggest PCSGs were poorly attended because men typically avoid disclosure due to low perceived need for support, fear of stigmatization, the need to minimize the threat of illness to aid coping, practical necessities in the work place, and the desire to avoid burdening others. Related to this, US-based and Australian-based men’s misperceptions that PCSG meetings were geared toward emotional support of the terminally ill (Krizek, Roberts, Ragan, Ferrara, & Lord, 1999; Walker, 2005) and a sense of shame and embarrassment about sharing personal information (Smith et al., 2002; Weber, Roberts, & McDougall, 2000) were identified as barriers to attending support groups.

There is also evidence that health care providers strongly influence men’s interest in attending a “support” group. For example, health care providers’ lack of awareness of PCSGs can be a barrier (Smith et al., 2002). A survey questionnaire study of 1,224 attendees at Australian-based PCSGs confirmed that the clinician’s level of endorsement for men’s participation in PCSGs strongly influenced them to attend (Steginga, Pinnock, Gardiner, Dunn, & Gardiner, 2005). This finding led researchers to conclude that clinicians are integral to the development and implementation of support services for men with PCa (Steginga et al., 2005). A subsequent study of 36 clinicians (27 urologists and 9 radiation oncologists) by Steginga et al. (2007) identified that participants were reluctant to refer patients to PCSGs, fearing that biased viewpoints and misinformation within the groups might contribute to men’s uncertainty and decisional regret.
In summary, there is consensus that PCSGs provide important resources to men and their families; however, group sustainability can be challenged by failing to attract new attendees, which in turn reduces the capacity for group leadership succession planning (Oliffe et al., 2008). Related to this it seems that men’s resistance to attending PCSGs might stem from concerns about how the groups operate and the legitimacy of the resources that they provide. In this regard, PCa specialists are well positioned to inform men about the availability of PCSGs. While PCa specialists’ reliability for reporting what may or may not motivate PCa patients to attend a support group is debatable, their collective views about PCSGs provide important insights to thoughtfully consider avenues for maximizing the potential of PCSGs.

**Subjects and Method**

This study was approved by the University of British Columbia’s Behavioural Research Ethics Board Committee. The survey instrument to solicit the PCa specialists’ views was developed as a web-survey composed of a 56-item questionnaire based on Stegenga et al.’s (2007) study tool. The survey items were first developed from in-depth expert stakeholder interviews with 36 clinicians (27 urologists and 9 radiation oncologists) who had provided care to men with PCa for a mean of 16 years. To confirm item relevance in an iterative approach a follow-up paper-based survey applying a 5-point Likert-type scale was completed by 30 of these participants from which key themes were identified. For the current study incorporated were five demographic questions and six sets of attitudinal items to measure beliefs about: positive influences of PCSGs, negative influences of PCSGs, reasons for attending PCSGs, the attributes of effective PCSGs, and the value of face-to-face and web-based PCSGs. Each set of attitudinal items included five to nine questions that were rated by respondents using a 5-point Likert-type scale (where 1 = strongly disagree and 5 = strongly agree). An open-ended question to provide additional information was also included. Reliabilities for the current study ranged from .72 to .94. This instrument is available on request.

In all, 150 PCa specialist respondents were recruited via existing professional contacts and the Canadian Urological Association. There was a nominal honorarium of a $50 gift card of their choice, which was mailed to respondents on receipt of the completed survey. Descriptive statistics are reported. Median scores are reported for ordinal Likert data. Responses to the open-ended questions were coded and content analyses performed to inductively derive broad descriptive categories. Respondents included 100 urologists, 40 radiation oncologists, and 10 medical oncologists. Table 1 presents descriptive data on the demographic and professional characteristics of the sample. Of 150 respondents, 82.7% (n = 124) were male and the majority of respondents practiced in British Columbia (38.0%) and Ontario (33.3%). Respondents worked in a variety of settings, including hospitals and/or cancer centers. They ranged in age from 26 to 86 years (mean age = 44.7 years) and had an average of 13.8 years’ experience working with PCa patients. The majority (62.0%) reported formally presenting information at PCSGs, and 78% indicated that they had referred patients to PCSGs.

**Results**

*Features of PCSGs That Positively Influence Men’s Adjustment to PCa*

Respondents provided ratings of seven features of PCSGs that were potentially positive influences on men’s adjustment to PCa (Figure 1). Ratings of these features were uniformly high (median of five). All seven characteristics (i.e., community support, reassurance, social identity, sharing experiences, emotional support, and providing information) were identified as important benefits. While friendship was rated the lowest of the items, sharing experiences and emotional support were highlighted as the strongest factors positively influencing men’s adjustment to PCa.

*Features of PCSGs That Negatively Influence Men’s Adjustment to PCa*

Specialists rated nine items related to possible negative influences of PCSGs including, meeting with dominant members who push their own views, dissemination of inaccurate information, hearing negative experiences, creating conflict over treatment decisions, promoting a specific clinician, creating confusion, supplying irrelevant information, causing confrontation, and promoting a single therapy (Figure 2). PCSG meetings with dominant members, dissemination of inaccurate information, and hearing negative experiences were identified as the three leading negative influences of PCSGs affecting men’s adjustment to PCa. The median response for the other six items was 3 (neutral) with no specific negative factors being identified as particularly influential. In addition, some negative influences were apparent in the open-ended responses discussed below.

*Reasons for Attending PCSGs*

The items most strongly endorsed by PCa specialists as the reasons that men attended PCSGs were to discuss PCa treatments and to gain information (median of 5; see Figure 3).
Reasons for Not Attending PCSGs

Perceived reasons for not attending PCSGs were assessed with seven items. Privacy issues was the most clearly endorsed reason for not attending PCSGs (median of 5), followed by ignorance of PCSGs (Figure 4). Items including “no need,” “having enough information,” and “a desire to move past the PCa experience,” were also highly endorsed by PCa specialists (median of 4) as reasons why some men might prefer not to attend PCSGs.

Characteristics of Effective PCSGs

Respondents rated items to assess their views on the characteristics of effective face-to-face PCSGs. Avoiding bias by not promoting one view of treatment, a range of different health care providers’ input, being patient-driven, and diversity of therapies discussed were the most highly endorsed characteristics of effective PCSGs (median rating of 5; Figure 5). Supported by other health care providers, having a trained facilitator, and support from health organizations were also generally agreed as keys to success (median rating of five).

Characteristics of Effective Web-Based PCSGs

Provision of summarized PCa information, the use of multimedia evidence-based health care provider presentations, and the facilitation of camaraderie were identified as the most essential features of effective web-based PCSGs (Figure 6). The findings suggest that specialists are most concerned with the accuracy and interactive capacity of the information made available online.

Additional Commentary

Thirty-four respondents shared additional insights via written comments. Table 2 presents the results from the content analysis of the open-text comments. In summary, the main cautions related to online PCSGs and prostate specific antigen advocacy. Online PCSGs were predicted to encounter similar shortcomings, and they were suggested to augment rather than replace face-to-face meetings for a variety of reasons including older men’s reduced access to online resources. In terms of endorsements and suggestions for promoting the work of PCSGs, respondents offered a range of ideas, including messaging individual clinicians about the groups and supplying brochures to clinics that patients could take with them.

Discussion

The results from the current study provide important insights to PCa specialists’ awareness of the benefits of PCSGs for men with regards to emotional support underpinned by the sharing of personal experience. Men are typically underrepresented as seekers of psychological support after cancer (Steginga et al., 2008) and a recent U.S. study indicated that PCa patients are approximately...
30% less likely than breast cancer patients to have discussed emotional and social concerns with their health care provider (Forsythe et al., 2013). In light of these findings it may be that although PCa specialists appear to understand and endorse the benefits of face-to-face PCSGs, emotional and social care, and therefore raising patient awareness about support groups is not necessarily a priority. That said, given that many men with PCa report unmet needs for support with regards to fears about the cancer returning or spreading and worry about “significant” others (Lintz et al., 2003; Smith et al., 2007; Steginga et al., 2001), PCSGs present as a valuable but potentially underused support program.

Contrasted within the current study findings that sharing experiences and emotional support were the most positive influences of PCSGs on men’s adjustment, PCa specialists rated information as men’s primary motivation for attending groups. Interestingly, the key reasons endorsed as potentially negative effects of support groups including online groups, centered on misinformation. This may reflect the uncertainty that persists about the optimal treatment approach for PCa, the expanding array of treatment options and regimes, and from this, clinicians concerns about supporting men’s decision making in the clinical encounter. Men with PCa often report high decisional conflict that for some persists over time (Steginga, Occhipinti, Gardiner, Heathcote, & Yaxley, 2004), and men do report others cancer experiences, both positive and negative, as influencing their treatment choices (Steginga et al., 2002). Hence, this is a highly salient clinical issue for specialists in these men’s care.

There was less agreement on the value of PCSG in terms of social interaction and helping other men with PCa, suggesting that some PCa specialists believe the
Figure 3. Reasons why men choose to attend a prostate cancer support group.

Figure 4. Reasons why men choose not to attend a prostate cancer support group.

Figure 5. Key factors for the effectiveness of an in-person prostate cancer support group.

The group format of PCSGs may be unappealing to men and dissuade them from attending. These views may also suggest that PCa specialists do not anticipate men to be long-term attendees.
Figure 6. Key factors for the effectiveness of a web-based prostate cancer support group.

Table 2. Main Endorsements and Cautions From Open-Ended Question.

| Endorsements                                                                 | Cautions                                                                 |
|----------------------------------------------------------------------------|--------------------------------------------------------------------------|
| • They have gradually transitioned over the years from consisting of those with bad experiences looking for support to those made up of men truly interested in helping their peers. | • Generally great people, but are really dependent on the skills of its members and limited by resources and contacts. I have had concerns in the past about messaging going out about prostate cancer screening from the support groups locally. |
| • They are important and should be available in all academic centers across Canada. | • A group with an open approach is best. Not focusing on any one treatment, or area. Avoid advocacy. |
| • Support groups are vital to the overall treatment of any disease particularly malignancies such as prostate cancer. | • I think face-to-face contact is important with web-based contact as an adjunct. |
| • Support groups of huge value to patients trying to make initial decision. | • Web-based solutions should link prostate cancer support groups across regions, to integrate smaller ones into bigger ones for enhanced support. |
| • I think it is worth making patients aware of so they can choose to be involved if it’s right for them. I think it is important that these types of things remain patient driven. | • A brochure that could be given out in clinic about support groups would be helpful—they could take home and have time to think about it and see what it’s about and reach out on their own rather than getting a yes or no in clinic if they are interested. |
| • Need more awareness of such programs. | • I have concerns about the quality and accessibility of web 2.0 options for support groups for many of our more elderly patients—I think they do however have potential for a strong role with our younger patients, families, and selected older patient populations. My fear having presented to these groups is that the range of disease is quite broad. |
| • Web-based solutions should link prostate cancer support groups across regions, to integrate smaller ones into bigger ones for enhanced support. | • In our area, the average participant at our support group is over 65, I am not sure they would participate to a web-based support group. |
| • A brochure that could be given out in clinic about support groups would be helpful—they could take home and have time to think about it and see what it’s about and reach out on their own rather than getting a yes or no in clinic if they are interested. | • I would think the benefits and goals of a web-based group may be different from an in-person group, and I’d have more concern about the web-based approach, regarding risks of misinformation, miscommunication, hostility from anonymous contributors, potential risks to confidentiality, etc. I wouldn’t personally feel that a web-based utility would be the optimal method for support group conduct or purposes. |
| • I’ve enjoyed participating educating and presenting at prostate cancer group. |                                                                 |
The overarching endorsement of specialists for face-to-face PCSGs suggest some shifts to Steginga et al. (2007) findings which highlighted specialists’ concerns about the potential for misinformation. Such subtle changes are likely influenced by an array of factors including specialists’ direct exposure to PCSGs, increased clinical emphasis on shared treatment decision-making, and the growth and research fundraising activities of groups more generally. While specialists’ concerns about misinformation may be more focused on web-based resources including online PCSGs, face-to-face groups may need to disect out and better define and focus their role in decision and information support if they are to further build specialist support and referral to their groups and programs.

Although this study is explorative, identified were some variations with respect to PCa specialist views about PCSGs. These findings as well as views that a lack of knowledge about PCSG might obscure PCSGs as a potential resource for men indicate that efforts are needed to raise awareness about PCSGs and its potential value in targeted and tailored communications to PCa specialists and men in the community. Engaging PCa survivors who have attended PCSGs to challenge assumptions and misconceptions about PCSGs that are barriers to attendance is likely to be helpful. The positive experiences of other men can provide powerful examples to overcome fears among potential attendees. However, a parallel educational initiative to ensure that PCa specialists are well informed about PCSGs is also needed. The consideration of PCa specialists’ personal perspective is essential in the planning and implementation of this type of initiative.

The current study limitations include recognition that the reliability of PCa specialists to report what may or may not motivate patients to attend a PCSG is debatable. Moreover, PCa specialists’ perspectives about PCSGs are one among many potentially diverse health care provider views. Therefore, acknowledged also are the limits in terms of generalizability in describing the viewpoints of a small Canada-based sample of a PCa specialists. Nonetheless, these limitations provide important direction for future studies, which might include large international multisite studies investigating qualitative interviews to more fully contextualize the findings offered in the current study.

While the PCSG sustainability issues raised by Oliffe et al. (2008) talked to the challenges connecting to nonprofit cancer agencies, the findings from the current study reveal the PCa specialists as allies with significant potential for advancing PCSGs, especially by raising awareness with potential attendees. In this regard, formalizing collaborations might be mutually beneficial whereby a steady flow of new PCSG attendees can be mustered and the burden on clinical psychosocial oncology services further reduced.

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References
Arrington, M. I. (2003). “I don’t want to be an artificial man”: Narrative reconstruction of sexuality among prostate cancer Survivors. Sexuality and Culture, 7(2), 30-58.
Arrington, M. I. (2010). Theorizing about social support and health communication in a prostate cancer support group. Journal of Psychosocial Oncology, 28, 260-268.
Arrington, M. I., Grant, C. H., & Vanderford, M. L. (2010). Man to man and side by side, they cope with prostate cancer: Self-help and social support. Journal of Psychosocial Oncology, 23, 81-102.
Australian Government. (2013). Prostate cancer statistics. Retrieved from http://canceraustralia.gov.au/affected-cancer/cancer-types/prostate-cancer/prostate-cancer-statistics
Botteroff, J. L., Oliffe, J. L., Halpin, M., Phillips, M., McLean, G., & Mroz, L. (2008). Women and prostate cancer support groups: The gender connect? Social Science & Medicine, 66, 1217-27.
Canadian Cancer Society. (2011). Canadian Cancer Statistics 2011: Featuring colorectal cancer. Toronto, Ontario, Canada: Canadian Cancer Society. Retrieved from http://www.cancer-asian.com/images/news/Canadian_Cancer%20Statistics_2011_English.pdf
Canadian Cancer Society. (2013). Prostate cancer statistics. Retrieved from http://www.cancer.ca/en/cancer-information/cancer-type/prostate-cancer/statistics?region=bc
Centers for Disease Control and Prevention. (2013). Prostate cancer statistics. Retrieved from http://www.cdc.gov/cancer/prostate/statistics
Chapple, A., & Ziebland, S. (2002). Prostate cancer: Embodied experience and perceptions of masculinity. Sociology of Health and Illness, 24, 820-841.
Charmaz, K. (1995). Identity, dilemmas of chronically ill men. In D. Sabo & D. F. Gordon (Eds.), Men’s health and illness: Gender, power and the body (pp. 266-291). Thousand Oaks, CA: Sage.
Cordova, M. J., Giese-Davis, J., Golant, M., Kronnenwetter, C., Chang, V., McFarlin, S., & Spiegel, D. (2003). Mood disturbance in community cancer support groups: The role of emotional suppression and fighting spirit. Journal of Psychosomatic Research, 55, 461-467.
Coreil, J., & Behal, R. (1999). Man to man prostate cancer support groups. Cancer Practice, 7, 122-129.
Fergus, K., Gray, R., & Fitch, M. (2002a). Active consideration: Conceptualizing patient-provided support for spouse caregivers in the context of prostate cancer. *Qualitative Health Research, 12*, 492-514.

Fergus, K., Gray, R., & Fitch, M. (2002b). Sexual dysfunction and the preservation of manhood: Experiences of men with prostate cancer. *Journal of Health Psychology, 7*(3), 303-316.

Forsythe, L. P., Kent, E. E., Weaver, K. E., Buchanan, N., Hawkins, N. A., Rodriguez, J. L., . . . Rowland, J. H. (2013). Receipt of psychosocial care among cancer survivors in the United States. *Journal of Clinical Oncology, 31*, 1961-1969.

Gannon, K., Guerro-Blanco, M., Patel, A., & Abel, P. (2010). Re-constructing masculinity following radical prostatectomy for prostate cancer. *Aging Male, 13*, 258-264.

Gray, R. E., Carroll, J. C., Fitch, M., Greenberg, M., Chart, P., & Orr, V. (1999). Cancer self-help groups and family physicians. *Cancer Research, 7*, 10-15.

Gray, R. E., Fitch, M. I., Davis, C., & Phillips, C. (1996). Breast cancer and prostate cancer self-help groups: reflections on differences. *Psycho-Oncology, 5*, 137-142.

Gray, R. E., Fitch, M. I., Davis, C., & Phillips, C. (1997). Interviews with men with prostate cancer about their self help group experience. *Journal of Palliative Care, 13*, 15-21.

Gray, R. E., Fitch, M., Phillips, C., Labrecque, M., & Fergus, K. (2000a). Managing the impact of illness: The experiences of men with prostate cancer and their spouses. *Journal of Health Psychology, 5*, 531-548.

Gray, R. E., Fitch, M., Phillips, C., Labrecque, M., & Fergus, K. (2000b). To tell or not to tell: Patterns of disclosure among men with prostate cancer. *Psycho-Oncology, 9*, 273-282.

Gregoire, I., Kalogeropoulos, D., & Corcos, J. (1997). The effectiveness of a professionally led support group for men with prostate cancer. *Urologic Nursing, 17*(2), 58-66.

Jemal, A., Siegel, R., Ward, E., Hao, Y., Xu, J., & Thun, M. J. (2009). Cancer statistics, 2009. *CA: A Cancer Journal for Clinicians, 39*, 225-249.

Katz, D., Koppie, T. M., Wu, D., Meng, M. V., Grossfeld, G. D., Sadesky, N., . . . Carroll, P. R. (2002). Socio-demographic characteristics and health related quality of life in men attending prostate cancer support groups. *Journal of Urology, 168*, 2092-2096.

Klemm, P., Bunnell, D., Cullen, M., Soneji, R., Gibbons, P., & Holecek, A. (2003). Online cancer support groups: A review of the research literature. *Computer Informatics Nursing, 21*, 136-142.

Klemm, P., Hurst, M., Dearholt, S. L., & Trone, S. R. (1999). Gender differences on internet cancer support groups. *Computers in Nursing, 17*(2), 65-72.

Krizek, C., Roberts, C., Ragan, R., Ferrara, J. J., & Lord, B. (1999). Gender and cancer support group participation. *Cancer Practice, 7*(2), 86-92.

Linz, K., Moyiniha, C., Steginga, S., Norman, A., Eeles, R., Huddart, R., . . . Watson, M. (2003). Prostate cancer patients’ support and psychological care needs: Survey from a non-surgical oncology clinic. *Psycho-Oncology, 12*, 769-783.

Manne, S. L. (2002). Prostate cancer support and advocacy groups: Their role for patients and family members. *Seminars in Urologic Oncology, 20*, 45-54.

Matsunaga, D. S., & Gotay, C. C. (2005). Characteristics contributing to an enduring prostate cancer support group in an Asian and Pacific Islander community. *Journal of Psychosocial Oncology, 22*(4), 1-30.

Meuleman, E. J. H., & Mulders, P. F. A. (2003). Erectile function after radical prostatectomy: A review. *European Urology, 43*, 95-102.

Navon, L., & Morag, A. (2003a). Advanced prostate cancer patients’ relationships with their spouses following hormonal therapy. *European Journal of Oncology Nursing, 7*(2), 72-81.

Navon, L., & Morag, A. (2003b). Advanced prostate cancer patients’ ways of coping with the hormonal therapy’s effect on body, sexuality, and spousal ties. *Qualitative Health Research, 13*, 1378-1392.

Navon, L., & Morag, A. (2004). Liminality as biographical disruption: Unclassifiability following hormonal therapy for advanced prostate cancer. *Social Science & Medicine, 58*, 2337-2347.

Oliffe, J. L. (2005). Constructions of masculinity following prostatectomy-induced impotence. *Social Science & Medicine, 60*, 2249-2259.

Oliffe, J. L. (2006). Embodied masculinity and androgen deprivation therapy. *Sociology of Health and Illness, 28*, 410-432.

Oliffe, J. L., Gerbrantd, J. S., Bottorff, J. L., & Hislop, T. G. (2010). Health promotion and illness demotion at prostate cancer support groups. *Health Promotion Practice, 11*, 562-571.

Oliffe, J. L., Halpin, M., Bottorff, J. L., Hislop, T. G., McKenzie, M., & Mroz, L. (2008). How prostate cancer support groups do and do not survive: British Columbian perspectives. *American Journal of Men’s Health, 2*, 143-155.

Remzi, M., Waldert, M., & Djavan, B. (2004). Prostate cancer in the ageing male. *American Journal of Men’s Health, 1*, 47-54.

Seale, C., Ziebland, S., & Charteris-Black, J. (2006). Gender, cancer experience and internet use: A comparative keyword analysis of interviews and online cancer support groups. *Social Science & Medicine, 62*, 2577-2590.

Smith, R. L., Crane, L. A., Byers, T., & Nelson-Marten, P. (2002). An evaluation of the man-to-man self-help group in Colorado and Utah. *Cancer Practice, 10*, 234-239.

Smith, D. P., Supramaniam, R., King, M. T., Ward, J., Berry, M., & Armstrong, B. K. (2007). Age, health, and education determine supportive care needs of men younger than 70 years with prostate cancer. *Journal of Clinical Oncology, 25*, 2560-2566.

Steginga, S. K., Campbell, A., Ferguson, M., Beeden, A., Walks, M., Cairns, W., & Dunn, J. (2008). Socio-demographic, psychosocial and attitudinal predictors of help seeking after cancer. *Psycho-Oncology, 17*, 997-1005.

Steginga, S. K., Oechipinti, S., Dunn, J., Gardiner, R. A., Healthcote, P., & Yaxley, J. (2001). The supportive care needs of men with prostate cancer. *Journal of Psychosocial Oncology, 10*(1), 66-75.
Steginga, S. K., Occhipinti, S., Gardiner, R. A., Heathcote, P., & Yaxley, J. (2002). Making decisions about treatment for localised prostate cancer. *British Journal of Urology International, 89*, 255-260.

Steginga, S. K., Occhipinti, S., Gardiner, R. A., Heathcote, P., & Yaxley, J. (2004). Prospective study of men’s psychological and decision-related adjustment after treatment for localized prostate cancer. *Urology, 63*, 751-756.

Steginga, S. K., Pinnock, C., Gardiner, M., Dunn, J., & Gardiner, R. A. (2005). Evaluating peer support for prostate cancer: The prostate cancer peer support inventory. *British Journal of Urology International, 95*, 46-50.

Steginga, S. K., Smith, D. P., Pinnock, C., Metcalfe, R., Gardiner, R. A., & Dunn, J. (2007). Clinicians’ attitudes to prostate cancer peer-support groups. *British Journal of Urology International, 99*, 68-71.

Thaxton, L., Emshoff, J. G., & Guessous, O. (2005). Prostate cancer support groups: A literature review. *Journal of Psychosocial Oncology, 23*, 25-40.

Voerman, B., Visser, A., Fischer, M., Garssen, B., Van Andel, G., & Bensing, J. (2007). Determinants of participation in social support groups for prostate cancer patients. *Psycho-Oncology, 16*, 1092-1099.

Walker, S. (2005). *Prostate cancer support groups: An evaluation* (Doctoral dissertation). Retrieved from http://research-bank.swinburne.edu.au/vital/access/services/Download/swin:7543/SOURCE2

Wall, D., & Kristjanson, L. (2005). Men, culture and hegemonic masculinity: Understanding the experience of prostate cancer. *Nursing Inquiry, 12*(2), 87-97.

Weber, B. A., Roberts, B. L., & McDougall, G. J. (2000). Exploring the efficacy of support groups for men with prostate cancer. *Geriatric Nursing, 21*, 250-253.

Weber, B. A., Roberts, B. L., Resnick, M., Deimling, G., Zauszniewski, J. A., Musil, C., & Yarandi, H. N. (2004). The effect of dyadic intervention on self-efficacy, social support, and depression for men with prostate cancer. *Journal of Psycho-Oncology, 13*(1), 47-60.

Weber, B. A., & Sherwill-Navarro, P. (2005). Psychosocial consequences of prostate cancer: 30 years of research. *Geriatric Nursing, 26*, 166-175.

Zanchetta, M. S., Cognet, M., Xenocostas, S., Aoki, D., & Talbot, Y. (2007). Prostate cancer among Canadian men: A transcultural representation. *International Journal of Men’s Health, 6*, 224-258.

Zhang, A. Y., Galanek, J., Strauss, G. J., & Siminof, L. A. (2008). What it would take for men to attend and benefit from support groups after prostatectomy for prostate cancer: A problem-solving approach. *Journal of Psycho-Oncology, 26*, 97-112.
