Men’s Information-Seeking Behaviour Regarding Cancer Risk and Screening:
A Meta-Narrative Systematic Review

Mohamad M. Saab*, Mary Reidy¹, Josephine Hegarty¹, Mairin O’Mahony¹, Mike Murphy²,
Christian Von Wagner³, Frances J. Drummond¹,₄

¹ School of Nursing and Midwifery, University College Cork, Ireland
² School of Applied Psychology, University College Cork, Ireland
³ Institute of Epidemiology and Health Care, University College London, United Kingdom
⁴ Department of Epidemiology and Public Health, University College Cork, Ireland

*Corresponding author at: Catherine McAuley School of Nursing and Midwifery, University College Cork, Brookfield Health Sciences Complex, College Road, Cork, Ireland.
Tel.: +353 21 490 1518
E-mail address: msaab@ucc.ie; mohamad.m.saab@gmail.com
ABSTRACT

Objective: Preventive strategies are known to reduce cancer risk and incidence and improve prognosis. Men seldom seek medical information about cancer prevention and risk reduction. The aim of this meta-narrative systematic review was to critically appraise evidence from qualitative, quantitative, and mixed-methods studies that explored men’s information-seeking behaviours in relation to cancer prevention and risk reduction.

Methods: MEDLINE, CINAHL Plus with Full Text, PsycINFO, PsycARTICLES, Psychology and Behavioral Sciences Collection, Education Full Text, and ERIC were systematically searched for studies published in English between January 1st 2006 and May 30th 2016. A total of 4,117 titles were identified; of which, 31 studies were included (21 qualitative studies, nine quantitative studies, and one mixed-methods study). The methodological quality of the studies was appraised using different tools.

Results: Most studies focused on screening for prostate (n=18) and colorectal cancer (n=7). The majority of men were passive information-gatherers rather than active information-seekers. Key sources of information included the internet for active information-seekers and healthcare professionals for passive information-gatherers. Barriers to information-seeking included information overload, embarrassment, and fear. Low literacy and health literacy levels were addressed in three studies and were identified as impediments to active information-seeking. Facilitators to information-seeking included family support, media, celebrity endorsements, and targeted information.

Conclusions: Men’s information-seeking behaviour regarding cancer risk reduction, prevention, and screening is influenced by several factors. This necessitates targeted interventions aimed at raising awareness of cancer prevention and screening, whilst accounting for men’s informational needs, preferred learning strategies, and literacy levels.

Keywords: cancer; colorectal cancer; health literacy; information-seeking; men; oncology; prevention; prostate cancer; screening; systematic review.
INTRODUCTION

Cancer remains one of the leading causes of mortality and morbidity around the world.¹ Fourteen million new cancer cases and 8.2 million cancer deaths were recorded worldwide in 2012.¹ It is expected that cancer incidence will increase by 70% over the coming two decades.¹ Cancer incidence and mortality are higher among men in comparison to women.² The most commonly diagnosed cancers in men are lung, prostate, colorectal, gastric, and liver cancer.¹²

Preventive strategies have been shown to reduce the risk of cancer and have been linked to improved prognosis.³ Nelson et al. reported a statistically significant decrease in breast cancer mortality in women aged 50 to 69 years who were screened compared to those who were not.⁴ Similarly, a randomised controlled trial with 30-year follow-up found that faecal occult blood testing reduced colorectal cancer mortality significantly when performed annually or biennially.⁵ While widespread, the effect of prostate cancer screening on mortality remains controversial.⁶,⁷ Other preventative strategies, including national smoking bans, have resulted in decreased smoking-related mortality.⁸ In order for preventative strategies to be effective, health information needs to reach, engage, and be understood by the target population.

In their analysis of the concept “health-seeking behaviour”, Lambert and Loiselle described health-seeking behaviours as “ways in which individuals go about obtaining information, including information about their health, health promotion activities, risks to one’s health, and illness” (p.1008).⁹ Health-seeking can be undertaken when people are asymptomatic, with a view to prevent disease.¹⁰ A range of information-seeking behaviours have been described, including active information-seeking (i.e. actively and with purpose seeking out information regarding a specific issue); active monitoring (i.e. actively scanning one’s environment for information, cues regarding a particular issue); passive monitoring (i.e. relying on chance encounters and other individuals to provide unsolicited information); and proxy searching (i.e. using intermediary channels such as friends or family members to search for information about an issue on behalf of the individual).¹¹

The literature reports that men seek health information less often than women.¹²,¹³ Moreover, men are more likely to engage in passive information-gathering, whereas women are more likely to be active information-seekers.¹³-¹⁵
Men’s lack of information-seeking is believed to be one of the leading causes for
deprioritising men’s health promotion among service providers and assuming that gendered
approaches to health should be primarily focused on women, rather than both genders
equally.\textsuperscript{16-18} This, according to Leone and Rovito, negatively impacts on men’s engagement
with health services and increases the health gap between both genders.\textsuperscript{19} In fact, men are less
likely to participate in preventative healthcare activities, including cancer screening, than
women,\textsuperscript{20} and are known to delay medical help-seeking for symptoms of male-specific,\textsuperscript{21} and
non-gender-specific malignancies.\textsuperscript{22}

In order to inform future health promotion policy and positively affect men’s health, it is
necessary to understand men’s health information-seeking behaviour in relation to cancer
prevention and risk reduction. To the authors’ knowledge, there has been only one systematic
review that focused on prostate cancer information.\textsuperscript{23} Therefore, the aim of this meta-
narrative systematic review was to critically appraise evidence from qualitative, quantitative,
and mixed-methods studies that explored men’s information-seeking behaviours in relation to
cancer prevention and risk reduction. This review was conducted based on five predefined
questions as follows:

(i) Where do men seek information on cancer prevention and risk reduction?
(ii) How do men use information on cancer prevention and risk reduction?
(iii) What are the barriers to information-seeking?
(iv) What are the facilitators to information-seeking?
(v) What is the impact of health literacy on information-seeking and use?

METHODS

This meta-narrative systematic review was conducted in accordance with the Cochrane
Handbook for Systematic Reviews,\textsuperscript{24} and reported using the 20-item Realist And MEta-
narrative Evidence Syntheses: Evolving Standards (RAMESES) reporting tool.\textsuperscript{25} Meta-
narrative review is a relatively recent systematic review methodology aimed at reviewing
evidence from qualitative and mixed-methods studies and is best suited for topics that have
been differently conceptualised.\textsuperscript{25} The review questions and methods were pre-defined and
were not changed during the review process.

Eligibility criteria
Empirical studies considered for inclusion met the following criteria: (i) involved men; (ii) adults (i.e. aged ≥18 years); (iii) primarily focused on where and how men seek and use information on cancer prevention and risk reduction; (iv) published between January 1st 2006 and May 30th 2016; and (v) published in English. Although there is no golden rule for limiting publications by date, scientific evidence published within a 10-year timeframe is considered to be recent.26,27 Eligible studies were included regardless of their methodological quality, since excluding studies on the basis of their methodological quality increases the risk of study selection and reporting bias.24

Studies involving women exclusively or where findings from men and women were indistinguishable were excluded. Studies involving cancer survivors were also excluded since their experiences may differ from those without cancer. Opinion papers, policy reports, and conference abstracts were excluded as their methodological quality could not be appraised. Dissertations and theses were excluded, since the merit of using them in systematic reviews is inconclusive.28

Information sources and search strategy

A systematic search of relevant electronic databases over a 10-year period was conducted. The electronic databases searched were: MEDLINE, CINAHL Plus with Full Text, PsycINFO, PsycARTICLES, Psychology and Behavioral Sciences Collection, Education Full Text, and ERIC. Each database was searched on May 30th 2016 for papers published between January 1st 2006 and May 30th 2016. Reference lists of eligible studies were checked for potentially relevant references that were not identified during the database search.

Boolean terms “OR” and “AND”, Medical Subject Headings (MeSH), and truncation “*” were used and the search was conducted on title or abstract as follows: (men OR males OR man OR male) AND (inform* OR advice OR advis* OR educat*) AND (cancer* OR neoplas* OR oncolog* OR tumour* OR tumor*) AND (need* OR necessit* OR require* OR seek* OR look* OR search* OR acquir* OR learn* OR "engag* with" OR use OR using OR utilis* OR utiliz*) AND (prevent* OR "reduce* risk" OR minimis* OR minimiz* OR "health promot*" OR screen*).

Study selection

Records identified through database searching were exported to Covidence, an online service recommended by Cochrane to facilitate data screening and extraction.29 Studies were first
screened on title and abstract to determine whether they met the inclusion criteria. The full-
texts of potentially relevant papers were then evaluated and reasons for excluding each article
were recorded. Title, abstract, and full-text screening were conducted independently by
paired reviewers. Disagreements were resolved by consensus and involved a third reviewer
when needed.

**Data collection process**

Data from included studies were extracted by all authors using a predefined extraction table.
Data were extracted as follows: author(s) and year; country and setting; study aim(s); study
design and theoretical underpinning; data collection method and instruments; and findings
relating to the review questions. The table was cross-checked for accuracy by the first author.

**Critical appraisal**

The choice of tools to appraise the methodological quality of the reviewed studies was
dependent on the study design. The 14-item Quality Assessment Tool for Observational
Cohort and Cross-Sectional Studies was used to critically appraise the quality of quantitative
studies. The overall quality of each study was rated as either ‘Poor’, ‘Fair’, or ‘Good’. The
quality of qualitative studies was appraised using the 10 items of the Critical Appraisal Skills
Programme (CASP) Qualitative Checklist. The Mixed Methods Appraisal Tool (MMAT)
comprising 13 questions in relation to the appropriateness of the qualitative methods,
quantitative methods, and the combination of both, was used to appraise the quality of mixed-
method studies. Each item in all three tools was evaluated on a ‘Yes’ and ‘No’ basis. Only
the Quality Assessment Tool for Observational Cohort and Cross-Sectional Studies enabled
an overall quality rating.

**Synthesis of results**

Data synthesis was conducted by the first author and cross-checked by the last author.
Findings were analysed and synthesised thematically according to the review aims.

Information-seeking behaviours were characterised as ‘active information-seeking’ and
‘passive information-gathering.’ Active information-seeking was defined as purposely
seeking out information and/or actively scanning one’s environment for information (i.e.
solicited information). Passive information-gathering was defined as relying on accidental
encounters and other individuals, including healthcare professionals, to provide unsolicited
information, and/or using intermediaries, such as friends and family members, to seek information on behalf of the individual.\textsuperscript{11}

Use of information to make a decision regarding cancer screening was investigated and factors influencing information-seeking were classified as barriers and facilitators. These pertained to information format, content, amount, and source.

Finally, the impact of literacy and health literacy on information-seeking and use was addressed. Health literacy was defined as the degree to which men were capable of obtaining, processing, and understanding information on cancer prevention and risk reduction.\textsuperscript{33}

**RESULTS**

**Study selection**

The study identification, screening, and selection processes are presented in Figure 1.\textsuperscript{34} Overall, 4,117 titles were identified through electronic database searching from MEDLINE (n=2,528); CINAHL Plus with Full Text (n=775); PsycINFO (n=670); PsycARTICLES (n=56); Psychology and Behavioral Sciences Collection (n=37); Education Full Text (n=28); and ERIC (n=23). Following deletion of duplicates, 3,374 records were screened on title and abstract and 3,054 irrelevant records were excluded. The full-texts of 320 papers were then evaluated and 289 articles were excluded. In total, 31 papers were deemed eligible for inclusion; 21 qualitative studies, nine quantitative studies, and one mixed-methods study. No additional studies were identified from reference list-checks.

**Study characteristics**

Study characteristics are presented in Table 1. The majority of the studies were conducted in the United States (n=20). Most of the participants were recruited from the community (n=12) and primary care practices and screening centres (n=7). Ten studies were underpinned by a theoretical framework. The minimum sample size for men was 8\textsuperscript{35,36} and the maximum was 4,194.\textsuperscript{37} Ages ranged between 19\textsuperscript{38} and 95 years.\textsuperscript{39} The majority of men were White (n=21). Studies focused primarily on information about prostate cancer and Prostate Specific Antigen (PSA) testing (n=18), followed by colorectal cancer (n=7).

**Critical appraisal**
The qualitative studies had clear aims and findings and used appropriate methods, designs, recruitment strategies, data analysis frameworks, and measures to enhance rigour. All but one qualitative study failed to address the relationship between the researcher and study participants, which increases the risk of bias (see supplementary Table 1S). The quantitative studies were rated as ‘Poor’ (n=4); ‘Good’ (n=3); and ‘Fair’ (n=2). Studies rated as ‘Poor’ failed to justify the sample size, specify the data collection timeframe, use valid and reliable data collection instruments, assess outcomes more than once, and/or adjust for confounders (see supplementary Table 2S). The mixed-methods study met all but one MMAT criterion, namely sample representativeness, since the quantitative element of this study was a pilot-test (see supplementary Table 3S).

Synthesis of results

For results from individual studies, see supplementary Table 4S.

Information-seeking behaviours and information sources

Men either sought information on cancer prevention and risk reduction themselves i.e. active information-seeking, or came across information i.e. passive information-gathering. In the majority of cases, the latter was true.

Active information-seeking

Men were predominantly active information-seekers in six studies: two qualitative and one quantitative longitudinal study assessing information-seeking for prostate cancer screening; one cross-sectional and one qualitative study on general cancer information-seeking; and one qualitative study on information-seeking for skin cancer prevention.

The internet served as the primary source of information among active seekers in five studies. Men searched for information on the internet to manage uncertainty following a high PSA test or to learn about the cancer diagnosis of a family member. Additionally, 61% of participants in a cross-sectional study actively sought general cancer information online. Haluza and Cervinka surveyed 193 men and 363 women about the impact of skin health information on their sun protective behaviours. It was found that women were more likely than men to seek information on skin cancer prevention using the internet (36.1% for men vs. 24.7% for women; p = 0.004). Other than the internet, 11% of participants in the longitudinal study by Gibson et al. actively sought PSA testing information from non-medical sources, including family, friends, and co-workers. Hicks et al. also
found that men’s social networks were identified as important sources of information, while participants in the study by Biddle et al. actively sought PSA testing information from “medical journals, newspapers, television, media, chat rooms, and medical brochures” (p.6) as well as their family, friends, and men who had similar experiences.

Passive information-gathering

Men passively acquired information in 13 studies: four quantitative, two qualitative, and one mixed-methods study on prostate cancer screening; three qualitative studies and one quantitative study on colorectal cancer screening; one quantitative study on Human Papilloma Virus (HPV) and anorectal cancer prevention; and one qualitative study on general cancer information.

Healthcare professionals were identified as a major source of unsolicited health information. For instance, participants in two qualitative studies identified their physician as the primary source of information about colorectal cancer. Blackwell et al. surveyed 89 men who have sex with men about their awareness of HPV and anorectal cancer. It was found that 49 participants had heard of anal Pap smears from different sources including their primary care physician (10.2%), nurse (16.3%), and other healthcare professionals (32.7%). This was also the case in five studies on prostate cancer prevention and PSA testing.

Men also identified numerous and varied nonmedical sources of passively acquired information including: friends, family members, and spouses; mass media; their churches; personal stories; and medical posters.

Use of information

Four studies addressed men’s use of acquired information to make informed decisions regarding cancer prevention and risk reduction practices. Of those, one was a qualitative study and one was a quantitative study on prostate cancer screening; one quantitative study addressed colorectal cancer screening; and one quantitative study explored general cancer information-seeking.

In a cross-sectional study aimed at exploring general cancer information-seeking in Australia, Zajac et al. found that men were less likely to search for information than women, but were more likely to be open to receiving unsolicited information (both p<0.001). However, only 32% of men expressed their willingness to receive unsolicited information via the internet.
Being equipped with information on cancer prevention and screening and having a close relative with a malignancy did not serve as predictors for prostate cancer screening.\(^\text{42}\) Similarly, a cross-sectional study examining the relationship between family support and PSA testing among men (n=625) found that those with a family member who had been diagnosed with prostate cancer were less likely to have undergone PSA testing in the previous year (odds ratio 0.47, 95% CI 0.31 to 0.70; p < 0.001).\(^\text{50}\)

Of note, prostate cancer screening remains controversial with conflicting evidence regarding the effectiveness of screening on reducing prostate cancer mortality.\(^\text{6,7}\) Furthermore, the opinions and screening behaviours of healthcare professionals differ, making the decision to undergo prostate cancer screening a difficult one for men.\(^\text{56}\)

As for colorectal cancer, “information-seeking by others was associated with greater absolute perceived risk of colon cancer, and information-seeking for oneself was associated with more frequent worry about colon cancer” (p.73).\(^\text{39}\)

**Barriers to information-seeking**

Barriers to information-seeking with regards to prostate cancer screening and PSA testing were addressed in eight qualitative,\(^\text{39,40,42,57-61}\) one quantitative,\(^\text{44}\) and one mixed-methods study.\(^\text{41}\) Barriers involved: information format and quantity (print and radio advertisements that are either lacking in or flooded with information ‘information overload’); information content (trigger words such as ‘research program’ and ‘research subjects’);\(^\text{38}\) lack of trust in the information offered by the media;\(^\text{57}\) and anxiety and fear.\(^\text{58}\)

Barriers to information-seeking were more pronounced among men belonging to some ethnic groups (i.e. Filipino, African American, and Latino men). In their qualitative study, Conde et al.\(^\text{40}\) found that first generation Filipino men were more likely to ignore prostate cancer information and Friedman et al.\(^\text{59}\) reported that African American men were primarily passive rather than active information-seekers. Moreover, two studies identified embarrassment, shame, perceived weakness, machoism, and fear as barriers to information-seeking in relation to prostate cancer among African American men.\(^\text{41,60}\) Lack of awareness of the PSA test, and cancer being a taboo subject among African American and Latino men also served as barriers to seeking information on prostate cancer and PSA testing.\(^\text{41,42,59,60}\) One study identified low literacy levels among African American men as a barrier to information-seeking.\(^\text{41}\) Furthermore, limited access to screening services/physicians and cost of screening were highlighted as barriers to information acquisition among African American men.\(^\text{41,59,61}\)
Similar barriers were reported in three qualitative studies on colorectal cancer. These include: lack of information and understanding, confusing instructions about faecal occult blood testing, generic materials being overlooked by certain cultures (e.g. American Indian); fear, embarrassment, clinic location/access, and concerns about privacy and cost.

**Facilitators to information-seeking**

Facilitators to information-seeking in relation to prostate cancer screening were addressed in nine qualitative, four quantitative, and one mixed-methods study. Many of these facilitators pertained to information layout, content, and mode of delivery. In two qualitative studies, men were more likely to acquire information using print media (e.g. sports section of the newspaper), appealing videos, and bullet points and favoured information that is practical and delivered via the mass media (e.g. television). Moreover, a number of men preferred information targeted towards men.

Men in a qualitative study who were asked to evaluate advertisements on prostate cancer screening, requested information about the signs, symptoms, and risk factors of prostate cancer and recommended using gender- and age-appropriate models and celebrities to promote screening. This was echoed in another study, whereby men interviewed about their prostate cancer information-seeking behaviour requested information on prostate cancer, its risk factors, and treatment options.

Community jurors (i.e. groups of men from the community) who evaluated different aspects of PSA testing believed that information about the risks and benefits of prostate biopsy and prostate cancer treatment should be offered to those who wanted it, including men with high PSA levels. They also believed that men would not want such information unless it was relevant to them.

Access to both, medical (e.g. trusted general practitioners and urologists) and non-medical (e.g. family and friends, the church, and neighbourhood settings) sources of information served as a facilitator to the passive acquisition of information.

Using information specifically targeted at different ethnic groups also facilitated information-seeking and acquisition. In two qualitative studies, African American men intended to seek information on ethnic-specific risk factors of prostate cancer, and recommended prostate cancer prevention messages that are simple, direct, and specific to African American males. Men also suggested that information should be delivered by trusted people including
African American church pastors, women, and prostate cancer survivors. Furthermore, transfer of knowledge between generations, being a second generation Filipino man who has computer-access, and living in a household that has access to information on prostate cancer (e.g. through newspapers), also served as facilitators to acquiring information on prostate cancer screening.

Four qualitative studies addressed facilitators to seeking colorectal cancer screening information. Bennett et al. found that factual information about colorectal cancer made screening less abstract. Moreover, men belonging to three different ethnic groups (i.e. African American, English Caribbean, and Haitian) recommended pamphlets at the doctor’s office, group sessions, and information and educational materials that are visual. Other facilitators included: using real people in narratives to make screening more vivid; leaflets with information about faecal occult blood testing; having a friend or a family member with cancer; and using social networks to raise awareness.

The impact of literacy levels on information-seeking and use

The impact of literacy and/or health literacy on information-seeking and use was addressed in two qualitative studies on colorectal cancer screening, and one mixed-methods study on prostate cancer screening.

Smith et al. explored, qualitatively, the colorectal cancer screening information needs and preferences of 14 men with varying literacy levels. Men with both high and low literacy levels reported that health information should be direct, short, and sharp and appreciated medical diagrams, found some of the statistics confusing and unclear, perceived some scientific information as difficult, and suggested phone helplines as a source of information. Moreover, both groups perceived medical terminology as problematic; however, this was more pronounced among the low literacy group. The lower literacy group also perceived high-density text as off-putting, was not reassured by scientific references, and perceived certain visual images as patronising. However, a weighing scale with ‘reasons to undergo colorectal cancer screening’ on one side and ‘reasons not to undergo colorectal cancer screening’ on the other side was perceived as helpful. Men with high literacy levels were reassured by scientific references and appreciated the use of visual images.

Friedman et al. conducted a mixed-methods study to assess functional health literacy among 25 African American men using two modified Cloze tests and the Shortened Test of Functional Health Literacy in Adults. Adequate comprehension of the survey tools was
demonstrated, with more than 56% correct answers. In addition, it was found that functional health literacy did not differ significantly by reading level and that 25% of participants were non-seekers of cancer information.\(^{41}\)

Finally, Smith et al. explored, qualitatively, how 12 men with low educational attainment used an evidence-based decisional aid to make colorectal cancer screening decisions.\(^{66}\) Understanding the purpose of the decisional aid determined how men used this information. Some used the information to make informed decisions on undergoing colorectal screening; some men chose to get screened because of the statistics; some chose to get screened despite doubting the statistics; and others chose not to get screened because they believed that the harms of screening outweighed its benefits. Moreover, some men dismissed this information and/or questioned its validity, including those who were critical of statistics and/or lacked the confidence to interpret statistics.\(^{66}\)

**DISCUSSION**

Findings from this meta-narrative systematic review suggest that men seek information in different ways, with the majority acquiring information passively through intermediary channels, rather than actively seeking this information. Men’s preferred format, content, and delivery of information were identified along with barriers and facilitators to seeking information on cancer prevention and risk reduction.

Regardless of how men acquire information; knowledge empowers.\(^{67}\) Therefore, once acquired, men can use information to make decisions about their health. In fact, the review found that men did use information to make decisions regarding cancer screening. However, very few studies explored the effect of literacy and/or health literacy on men’s understanding of cancer prevention and risk reduction information, which could negatively impact decision-making.\(^{41,65,66}\)

Knowledge of how and where men acquire information is required to ensure its effectiveness. An individual’s information field is the totality of possible sources an individual may consult and incorporates their information network. The daily sphere of information, i.e. the source of information an individual comes into contact with on a daily basis, is most likely to be with those whom they perceive that they have shared interests, for example, sports teams/groups, work associates, peer support groups, religious groups, friends, and neighbours.\(^{44,68,69}\)
Additionally, this can include journals, newspapers, television, internet, chat rooms, medical brochures, and speaking to family members, friends, and men who have similar experiences. As an exemplar, a large proportion of men read the sports sections of newspapers on a daily basis; understanding this is very important as the daily sphere of information varies for each man and differs according to their age, cultural, and socioeconomic backgrounds. Adolescent and young adults regularly name their parents, peers, and teachers as their key social network and the internet and social media as the key place for accessing information. In contrast, healthcare professionals are regularly cited as a major source of information for older males.

African American men have preferences for messages delivered through word of mouth and from credible sources such as African American church pastors, women, and prostate cancer survivors. Thus, for targeted health promotion interventions, it is useful to consider the target audience, their social networks, and their likely daily sphere of information. In addition, the informational preferences of the target group are important.

This review revealed that men preferred gender- and age-specific information presented as practical, factual, simple, and direct information using bullet points, pictures/models and multimedia. Moreover, endorsement of information by others was a feature mentioned in a number of studies; including endorsements by celebrities and receiving information through trusted sources such as religious figures.

Findings from this review are echoed in the wider literature on health information-seeking. In a study aimed at identifying the demographic characteristics of health information-seekers, Kim found that inactive information-seekers were predominantly males. Similarly, a survey aimed at determining the characteristics of online health information-seekers, found that men were less likely than women to engage in active health information-seeking using the internet.

One explanation for the disparity between the two genders, is that health-seeking is often perceived as a ‘feminine’ rather than ‘masculine’ behaviour. This could be explained further using sociological theories, including Connell's Theory of Hegemonic Masculinity which describes how masculine social constructs, such as stoicism, risk-taking, and toughness, impinge on the ways men seek health information. Men need to be able to justify engaging with healthy lifestyle behaviours and health services.
In the present review, men belonging to different ethnic groups (i.e. Filipino, African American, and Latino) were predominantly passive information-gatherers. Similar findings were identified in the wider literature on health information-seeking, whereby Latino and African American men identified their healthcare providers as the primary source of unsolicited health information. Age and health literacy were also found to impact on wider health information-seeking. For instance, in a study exploring internet use among low-income adults, Jensen et al. found that older individuals were less likely to seek health information online, mainly due to low levels of health and computer literacy.

**Implications for future research**

The present review has a number of research implications. For men to engage with cancer prevention information, they must perceive it as relevant to them; thus the content and information transmission processes need to be nuanced to reflect gender, generational, and ethnic differences. Moreover, information needs to be developed in both gender-specific and gender neutral formats where appropriate, as it has been shown that gender-specific information can have adverse effects on a number behaviours related to cancer prevention and information-gathering among women. It is also essential that the target audience, preferred learning strategies, social networks, and daily sphere of information are considered when designing health promoting interventions. While older men might benefit from simple interventions that do not involve complex technologies, interactive and visually appealing interventions can be used successfully to target younger men. In addition, understanding the age profile of cancers is important, with prostate and colon cancer information being more relevant for older men and testicular cancer information being more relevant to younger men. However, educational interventions aimed at schools are likely to have more lifelong implications on men’s health.

Researchers need to be vigilant for the potential of information overload and low literacy levels and are encouraged to design interventions that are non-patronising, yet easy to understand. Future research is also required to understand the cognitions and behaviours of passive information-seekers using theories such as the Cognitive Information Processing Theory. This in turn will inform the development and testing of targeted interventions to increase information-seeking among passive information-gatherers and help them to readily encode, store, and retrieve information.
The internet was identified as the key source of health information among active information-seekers. Consequently, evidence-based information needs to be developed and updated periodically using trustworthy online platforms.

From a methodological perspective, designing interventions that are underpinned by theory and using valid and reliable data collection instruments is required to strengthen the evidence base. An example is the M.A.L.E. H.E.L.P. questionnaire developed by Leone et al. to assess men’s knowledge, attitudes, and behaviours in relation to access to health care. Finally, longitudinal research is needed to explore the impact of information-seeking on cancer risk, incidence, and mortality.

**Implications for clinical practice**

Healthcare professionals, including nurses and physicians, were identified as the main source of unsolicited health information among passive information-gatherers. Therefore, increasing the participation of healthcare professionals in targeted health promoting activities including workshops, conferences, and continuing education may increase information acquisition among their patients and the wider community.

Healthcare professionals need to be cognisant of men’s sociodemographic characteristics including age, ethnicity, educational attainment, and level of health literacy, whilst providing information and advice in relation to cancer prevention and screening. These could be in the form of pamphlets, brochures, and/or posters placed in waiting areas and/or websites of national and international cancer organisations that offer easy to understand patient information.

Health organisations, where appropriate, are encouraged to adopt non-gendered approaches, whereby both men and women have equal opportunities to access care. This could be achieved by addressing structural barriers such as screening locations and times; and factors such as socioeconomic status, ethnicity, age group, and level of literacy and health literacy in cancer prevention information.

**Limitations**

The reviewed literature on men’s information-seeking behaviour in relation to cancer risk and screening was limited to a few cancers – being dominated by prostate and colorectal cancer screening. It is estimated that one third of cancer cases could be reduced through adherence to
the recommendations in the European Code Against Cancer; therefore, there is a need to expand the sphere of research on men’s health information seeking and engagement. Understanding how to engage men with this information remains largely unexplored. In addition, the reviewed studies were primarily conducted in United States, and barriers and facilitators in that health system may not be generalizable to men in other health systems. Furthermore, despite low health literacy being so prevalent, the impact of low health literacy on health information-seeking has not been well researched. Methodologically, all but one qualitative study failed to address the relationship between the researchers and participants and the quality of four of the nine quantitative studies was rated as ‘Poor’, which increases the potential for bias.

A number of limitations at the review level are also noteworthy. The search was limited to seven electronic databases, did not include records from the Grey literature, excluded theses and dissertations, and only included studies published in English between January 1st 2006 and May 30th 2016, which increases the risk of study selection bias. Moreover, only studies that are in line with the review aim and questions were included, which leaves room for reporting bias.

**CONCLUSION**

Findings from the present review highlight the need to: (i) explore the behaviours underlying passive information-gathering among men; (ii) empower men who are passive information-gatherers to engage with cancer prevention and risk reduction information; (iii) provide men with trustworthy and accessible information platforms; (iv) encourage healthcare professionals to partake in targeted health promoting activities; (v) use men’s daily sphere of information to increase awareness of cancer prevention; (vi) explore men’s information needs and preferred learning strategies; and (vii) design and measure the impact of targeted interventions aimed at men of different ages, socio-economic and ethnic groups, and literacy and health literacy levels.

**Funding:** This systematic review was undertaken as part of the MeCHanic study which was funded by the Irish Cancer Society grant HEA16DRU.

**Conflict of interest:** The authors declare no conflicts of interest.
References


17. Lefkowitch M, Richardson N, Robertson S. “If we want to get men in, then we need to ask men what they want”: pathways to effective health programing for men. *Am J Mens Health.* doi:10.1177/1557988315617825


<table>
<thead>
<tr>
<th><strong>Table 1. Study characteristics (n=31)</strong></th>
</tr>
</thead>
</table>
| **Country** | USA (n=20)  
Australia (n=5)  
UK (n=5)  
Dominican Republic (n=1) |
| **Setting** | Community (n=12)  
Health centres/clinics (n=7)  
Colleges (n=4)  
Pre-existing databases (n=3)  
Churches (n=2)  
State service agencies (n=2)  
Social event (n=1) |
| **Study design** | Qualitative:  
Descriptive (n=15)  
Exploratory (n=2)  
Grounded theory (n=2)  
Community jury (n=1)  
Descriptive and exploratory (n=1)  
Quantitative:  
Cross-sectional (n=5)  
Descriptive (n=1)  
Longitudinal (n=1)  
Needs assessment (n=1)  
Retrospective (n=1)  
Mixed-methods (n=1) |
| **Theoretical underpinning** | Community Based Participatory Model (n=2)  
Andersen’s Behaviour Model of Health Services (n=1)  
Health Belief Model (n=1)  
Nutbeam’s Health Literacy Framework (n=1)  
Preventive Health Model (n=1)  
Social Support Theoretical Framework (n=1)  
Systemic Linguistic Theory (n=1)  
Theory of Planned Behaviour (n=1)  
Theory of Reasoned Action (n=1)  
None/Not reported (n=21) |
| **Sample size (min-max)** | 8–4,194 |
| **Ethnic groups** | White (n=21)  
Black (n=7)  
American Indian (n=1)  
Filipino (n=1)  
Hispanic (n=1) |
| **Primary focus** | Prostate cancer (n=18)  
Colorectal cancer (n=7)  
General cancer information (n=4)  
Skin cancer (n=1)  
HPV/anorectal cancer (n=1) |