Prevalence and alternative explanations influence cancer diagnosis: an experimental study with physicians

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Abstract

Objective: Cancer causes death to millions of people worldwide. Early detection of cancer in primary care may enhance patients’ chances of survival. Physicians, however, often miss early cancers, which tend to present with undifferentiated symptoms. Within a theoretical framework of the hypothesis generation model (HyGene), and psychological literature, we studied how two factors – cancer prevalence and an alternative explanation for the patient’s symptoms – impede early cancer detection, as well as prompt patient management. Methods: Three hundred family physicians diagnosed and managed two patient cases, where cancer was a possible diagnosis (one colorectal, the other lung cancer). We employed a 2 (cancer prevalence: low vs. high) × 2 (alternative explanation: present vs. absent) between-subjects design. Cancer prevalence was manipulated by changing either patient age or sex; the alternative explanation for the symptoms was manipulated by adding or removing a relevant clinical history. Each patient consulted twice. Results: In a series of random-intercept logistic models, both higher prevalence (OR 1.92, 95% CI [1.27, 2.92]) and absence of an alternative explanation (OR 1.70, 95% CI[1.11, 2.59]) increased the likelihood of a cancer diagnosis which, in turn, increased the likelihood of prompt referral (OR 22.84, 95% CI[16.14, 32.32]). Conclusions: These findings confirm the probabilistic nature of the diagnosis generation process and validate the application of the hypothesis generation model to early cancer detection. Increasing the salience of cancer – such as listing cancer as a diagnostic possibility – during the initial hypothesis generation phase may improve early cancer detection.

Keywords: cancer diagnosis, prevalence, alternative explanation, diagnostic reasoning, diagnostic support
Cancer is one of the most common causes of mortality in the world (Stewart & Wild, 2014). It is also a disease that often gets misdiagnosed (Kern, 1994; Singh et al., 2013). Since early detection of cancer increases the chances of successful treatment, minimizing diagnostic delays is crucial (Møller et al., 2015; Neal, 2009; Richards, 2009). Substantial variation in urgent cancer referrals in UK primary care has caused concern (Baughan, Keatins, & O’Neill, 2011; Meechan et al., 2012). Physician-related cognitive factors, such as failure to consider cancer as a possible diagnosis, might account for some of this variation (Richards, 2009). Therefore, the present paper focuses on studying some of the cognitive factors affecting cancer generation and the evaluation processes used by family physicians via an experimental vignette study.

Cognitive factors are a common cause of diagnostic errors (Graber, Franklin, & Gordon, 2005; Norman & Eva, 2010). Retrospective reviews of litigation cases and patient charts have inferred that cognitive factors are the commonest contributors to diagnostic error (Gandhi et al., 2006; Lyratzopoulos, Vedsted, & Singh, 2015). Psychological studies have indeed identified several cognitive factors that can influence the outcome of the diagnostic process: hypothesis generation (e.g., Kostopoulou, Mousoulis, & Delaney, 2009; Kostopoulou, Siroti, Round, Samaranayaka, & Delaney, 2016); information search and interpretation driven by the leading hypothesis (Kostopoulou et al., 2009; Nurek, Kostopoulou, & Hagmayer, 2014); and primacy effects (Chapman, Bergus, & Elstein, 1996; Rebitschek, Bocklisch, Scholz, Krems, & Jahn, 2015). Research therefore strongly indicates that having an appropriate hypothesis – which could be determined by the initial cues perceived, and which determines subsequent information search and interpretation – plays a crucial role in the diagnostic process.

Accounts from the clinical reasoning literature discuss hypothesis generation as matching of the presenting problem to a similar problem previously encountered (Norman & Brooks,
1997), abstraction from specific patient features to higher-order memory structures (Grant & Marsden, 1987), and mapping specific patient cases to disease prototypes (Charlin, Tardif, & Boshuizen, 2000). For a review, see Kostopoulou (2009). Accounts from the psychological literature discuss the hypothesis generation as an employment of preconscious heuristic processes, which form the most plausible epistemic mental models of the world (Evans, 2006, 2008; Evans, Venn, & Feeney, 2002). These accounts differ in the knowledge or memory structures that they posit but are similar in that they underspecify the hypothesis generation process.

In contrast to previous theoretical accounts, the Hypothesis Generation (HyGene) model is a computational process model of hypothesis generation that attempts to specify this process more precisely (e.g., Thomas, Dougherty, & Buttaccio, 2014; Thomas, Dougherty, Sprenger, & Harbison, 2008). The model assumes three basic cognitive processes: 1/ retrieval from memory, 2/ maintenance in working memory, and 3/ judgment and decision-making. Specifically, the patients’ presented symptoms activate traces in episodic memory, representing similar past cases. An “unspecified probe” extracts the hypotheses (diagnoses) that are most frequently and strongly associated with the symptoms, and matches them against known hypotheses (e.g., prototypes of diseases stored in semantic memory). Hence, the better the match between cancer prototype and the patients’ symptoms, the more likely it is that the cancer hypothesis will reach the minimum activation threshold to enter working memory. This is an iterative process, so once an additional hypothesis is sufficiently activated by the probe (i.e., achieves the dynamically updated minimum activation threshold), it too enters working memory. However, due to memory constraints, physicians would only keep a limited number of hypotheses in their working memory; these would then guide subsequent information search and hypothesis testing. A
process comparing the relative strengths of the competing hypotheses in working memory
determines the posterior probability of the hypotheses. A hypothesis-guided search starts and
proceeds if more than one hypothesis is tested. The model assumes a consistency check of the
hypothesis with the input data: a hypothesis is rejected if it is not consistent with the symptoms.

Several factors may affect the generation of cancer hypotheses (e.g., time constraints and
working memory constraints) but two factors are especially likely to determine the degree of a
match between patient symptoms and the cancer hypothesis, and therefore its strength of
activation and maintenance in working memory: the a priori probability of cancer and the
presence of another explanation (a more common disease that also matches the patients’
symptoms). First, physicians perceiving a low probability of cancer in patients presenting with
vague symptoms may fail to generate the cancer hypothesis or exclude it as an implausible
explanation for the patients’ symptoms in the hypothesis evaluation phase. Indeed, people tend to
first generate the hypotheses with a high a priori probability (Dougherty & Hunter, 2003a;
Weber, Böckenholt, Hilton, & Wallace, 1993). Physicians may also fail to diagnose cancer if
there is an alternative, more common, explanation for the symptoms in an early stage of the
diagnostic process. This might impair either the hypothesis generation process – the cancer
diagnosis does not reach the dynamically updated activation threshold (Gettys & Fisher, 1979) –
and/or the hypothesis evaluation process – the relative strength (and hence the posterior
probability) of the cancer hypothesis is weakened due to a stronger plausible alternative
(Dougherty & Hunter, 2003a; Windschitl & Wells, 1998).
Present research

In the experiment presented here, we tested the effect of prevalence and an alternative explanation on the diagnosis and subsequent management of early cancers. We constructed two patient cases that depicted common cancers: one colorectal and the other lung. Colorectal and lung cancers are some of the most common cancers (Maddams, Utley, & Möller, 2012), and are often missed in their early stages of presentation (Rubin, McPhail, & Elliott, 2011). We avoided well-known methodological pitfalls. First, we manipulated the two factors using a factorial design, while keeping confounding variables, such as disease severity, constant. Second, we designed the clinical vignettes and manipulated the study factors in an ecologically valid way: i) we manipulated patient characteristics to alter disease prevalence rather than provide descriptive base rates, and ii) we either provided or withheld a medical history that could offer a plausible, alternative explanation for the patients’ symptoms. Finally, to overcome possible pitfalls of novice-experts differences (e.g., Shanteau, 1992), we recruited practicing physicians, rather than medical students, as study participants. We employed an experimental vignette study because it offers high internal and external validity of clinical judgments (Evans et al., 2015; Rose et al., 2015).

We tested three hypotheses. First, we hypothesized that a set of symptoms in patients with a higher prevalence of cancer will lead physicians to diagnose cancer more often than with a lower prevalence (Hypothesis 1). Second, we hypothesized that a set of symptoms in patients with a prior, alternative (i.e., non-cancerous) explanation of their symptoms will lead physicians to diagnose cancer less often than when an alternative explanation is not available (Hypothesis 2). Finally, as shown in previous research (e.g., Kostopoulou et al., 2008), we expected that diagnostic decisions would affect management decisions (Hypothesis 3).
Method

Participants

We determined our sample size in two steps. First, we defined the sample size stopping rule based on the a priori power analysis using G*Power (Cohen, 1988; Faul, Erdfelder, Lang, & Buchner, 2007). We required a minimum sample size of 237 participants for a logistic regression model with a binary predictor and adjusted for a covariate accounting for an additional 20% – to detect an odds ratio of 2.3, with 80% power at a 5% significance level. Such an effect size was assumed in other studies that used hypothetical vignettes to study the effect of patient demographics on diagnostic accuracy (Eva, Link, Lutfey, & McKinlay, 2010). Second, we increased the computed sample size by around 25% (i.e., n = 297) to account for the number of participants who would refer at least one patient on the first visit.

We invited, by e-mail, fully certified and currently practicing family physicians, who had participated in previous studies conducted by the lab. We also received help with recruitment from local clinical research networks around the UK, who contacted primary care clinics on our behalf. Potential participants were told that they would be taking part in a study of clinical reasoning. Cancer was not mentioned in the study information. After completing the study, each respondent was asked to provide an e-mail address in order to receive a £20 electronic Amazon voucher. To ensure that participants were actually physicians, and that each participant completed the study only once, we asked them to provide their professional NHS (National Health Service) e-mail address. Participants’ e-mail addresses were not linked to their data.

We recorded 382 attempts to complete the questionnaire. Only participants who completed all the vignettes were entered into the analysis. We excluded 82 incomplete attempts, most of which only explore the first presented vignette. As a result, 300 family physicians
completed the questionnaire. Half of the sample (50.7%) were male and 36% of participants were currently practicing in an inner city, while 45.7% practiced in an urban area and 18.3% in a rural area. Experience in family medicine post-certification ranged from 0 to 40 years ($M = 10.7$, $SD = 9.2$ years, $Mdn = 7$).

**Design and Materials**

We employed a 2 (cancer prevalence: low vs. high) × 2 (alternative explanation: present vs. absent) between-subjects experimental design for both patient cases (colorectal cancer and lung cancer), respectively. Each patient consulted twice. We therefore constructed two vignettes per case; one per consultation. We constructed the vignettes with the help of two experienced family physicians (the last two co-authors), who systematically reviewed the clinical literature on predictive symptoms and signs of colorectal and lung cancers. Participants saw both vignettes per case, unless they decided to refer the patient at the first consultation. In the colorectal cancer case, we manipulated cancer prevalence by changing the patient’s age (40 vs. 70 years for low vs. high prevalence respectively) to reflect increasing incidence with age (e.g., the increased incidence for UK males aged over 50, Cancer Research UK, 2012a). In the lung cancer case, we manipulated cancer prevalence by changing the patient’s sex to reflect the past trend of higher prevalence of lung cancer in men vs. women (e.g., age-standardised incidence rates in the UK are higher for males compared with females: 95.7 per 100,000 males and 66.5 per 100,000 females, Cancer Research UK, 2012b). The alternative explanation in the colorectal cancer case was hemorrhoids; in the lung cancer case it was infective exacerbation of pre-existing COPD (chronic obstructive pulmonary disease). The alternative explanation was either present or absent. Thus, we created four vignette versions per consultation and patient case. The complete set of vignettes and their versions is presented in the Appendix.
Second consultation vignettes always followed first consultation vignettes. At the first consultation, participants were randomized to either the colorectal or the lung cancer case. They saw and responded to the first consultations of both cancer cases. At the second consultation, they were randomized again to either the colorectal or the lung cancer case. Between these second consultations they saw a single-consultation vignette, which was not about cancer (a child presenting with an ear infection). This served as a decoy, to minimize any suggestion that the study was about cancer, and was also used to test a different hypothesis, not related to this study. No data from the decoy case are presented here.

Each case included basic information about the patient (name, sex, age, smoking status, last recorded blood pressure, relevant past medical history, current medications, and the reason for the last consultation), the current health complaint, current symptoms, and the results of relevant physical examinations and investigations. At the second consultation, the patients were still suffering from the same or slightly worsened symptoms, and had some new symptoms; specifically, the colorectal cancer patient presented with two “alarm” symptoms (blood in stool and slightly reduced hemoglobin indicating possible anemia and therefore blood loss), while the lung cancer patient presented with additional vague symptoms (fatigue, loss of appetite), which are not cancer-specific.

**Procedure**

Once participants had accessed the study site and read some information about the study, they were asked to provide socio-demographic details: gender, number of years in family medicine and location of their practice (inner-city, urban, or rural). They then read a vignette about the first consultation of the first patient (either colorectal or lung cancer). At the end of the vignette, they confirmed that they had read it, before proceeding to the diagnosis and
management page. First, they were asked to enter their main diagnosis ("What is your working diagnosis? (Please enter only one)"). They could also enter any other diagnoses that they were considering (see Supplementary materials). They were then asked to select one or more of the following list of management options: “arrange follow-up”, “order investigations”, “prescribe”, and “refer to secondary care”. Some of these options could be specified further. For example, the option “refer to secondary care” required a reason for referral, the type of specialist and the type of referral: emergency (within 24 hours), urgent (within two weeks – as recommended for suspected cancer) or routine (typically four to eight weeks). The management options fully respected the UK healthcare system to make the diagnostic and management task as realistic as possible. The guidelines for urgent referral for suspected cancer are laid down by NICE (NICE, 2015). At the end of the study, participants entered their e-mail address in order to receive the Amazon voucher.

We created two diagnostic variables: i) cancer as the main diagnosis: whether cancer was the main diagnosis or not and ii) cancer as the diagnosis: whether cancer was recorded at all, either as the main diagnosis or as one of the other diagnoses considered (the “differential”). The first author coded whether participants provided a cancer diagnosis or not for both diagnostic variables on each occasion. One of the physician co-authors coded whether the management decisions were likely to lead to early, delayed or nil detection of cancer (e.g., a decision not to refer was coded as not leading to cancer detection; a decision to refer to the wrong specialty was coded as leading to delayed cancer detection). The other physician co-author coded the management decisions on a subset of responses ($n = 657$, 57.6%). Inter-rater agreement was very high: 98% agreement (Cohen’s $\kappa = 0.96$, SE = 0.03) and the few disagreements were resolved in the discussion. We recoded the management decisions in two ways, one as early cancer
management (early cancer referral vs. late/no cancer referral) and one as cancer management (any cancer referral vs. no cancer referral).

We conducted the study in accordance with the ethical standards of the American Psychological Association, and obtained ethical approval from the Proportionate Review Sub-Committee of West London (REC 2).

**Results**

In total, the physicians considered cancer 70.2% of the time (800 out of 1,140 responses), while cancer was their main working diagnosis only 23.3% of the time (266 out of 1,140 responses). They did not refer for cancer at all 65.7% of the time, referred for cancer routinely 10.6% of the time, and urgently 23.7% of the time. Thus, although most responses included cancer as a diagnostic possibility, they did not always include a referral decision. The physicians recorded cancer as their main diagnosis more frequently in the colorectal cancer case than in the lung cancer case, and at the second rather than the first consultation (Figure 1, panel A). A similar pattern was seen in the urgent cancer referrals (Figure 2, panel A). In relation to our manipulated factors, the physicians recorded cancer as their main diagnosis more often in the vignettes where cancer prevalence was high (vs. low) and in the vignettes where there was no prior alternative explanation for the symptoms (vs. an alternative explanation) (Figure 1, panel B). Again, diagnostic rates were reflected in the urgent referral rates (Figure 2, panel B).

We measured the influence of cancer prevalence (Hypothesis 1) and an alternative explanation (Hypothesis 2) on diagnosis, using generalized estimating equations (GEE) – logistic
regression with random intercept, and vignette sequence as a repeated measure. In a model without covariates (Table 1, left column), higher cancer prevalence and absence of an alternative explanation almost doubled the frequency of cancer as the main diagnosis. There were no significant interactions between the two factors. In a model adjusted for covariates (Table 1, left column) – consultation (first and second) and physician experience (years in family medicine) – produced similar results: higher prevalence and absence of an alternative explanation increased cancer diagnosis, without a significant interaction. The physicians diagnosed cancer more than four times more often at the second than at the first consultation, and more experienced family physicians diagnosed cancer slightly more often than less experienced physicians. When we repeated these analyses for the more inclusive measure of diagnosis (cancer given as either the main diagnosis or in the differential, Table 1, right column), only prevalence remained significant. In terms of the covariates (Table 1, right column), consultation decreased and physician experience increased the odds of diagnosing cancer. Therefore, when consultation and physician experience were accounted for in the model, prevalence increased the odds of recording cancer, either as the main diagnosis or in the differential, whereas the alternative explanation increased the odds of cancer recorded as the main diagnosis only. Such evidence provides support for the two hypotheses derived from the hypothesis generation model about the effect of prevalence (Hypothesis 1) and an alternative explanation (Hypothesis 2) on cancer diagnosis.

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Insert Table 1 around here
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We had expected that, if cancer was the participants’ main diagnosis, this would lead to
Early detection of cancer (Hypothesis 3). Indeed, a series of GEE logistic regression models with random intercept whether unadjusted or adjusted for consultation and physician experience, confirmed this expectation (Table 2). Recording cancer as the main diagnosis increased the odds of any cancer referral (vs. no referral) around 13 times in both unadjusted and adjusted models, and the odds of early cancer referral (vs. delayed or no referral) almost 23 times in the unadjusted and 20 in the adjusted model. A second consultation also increased the odds of both referral (vs. no referral) and urgent referral (vs. delayed or no referral), on average, by a factor of 5. Physician experience was not related to referral decisions (Table 2). Thus, Hypothesis 3 was also confirmed: having cancer as the main diagnosis led to a higher rate of referrals.

Discussion

The goal of this study was to investigate two possible determinants of the prompt detection and management of early cancers by family physicians: disease prevalence and the availability of a prior, alternative explanation. Specifically, higher prevalence and no alternative explanation for the symptoms increased the frequency of cancer diagnoses and referrals to secondary care. In addition, we found that physicians gave cancer as their main working diagnosis and referred the patient urgently (i.e., within two weeks) much more often with repeated presentations that suggested a worsening of the condition, and more often for the colorectal cancer case, which included “alarm” symptoms, than for the lung cancer case, which included only vague symptoms.

Physicians’ sensitivity to prevalence is consistent with the hypothesis generation model.
and the literature in the field (Dougherty & Hunter, 2003a; Thomas et al., 2008; Weber et al., 1993). It appears to contrast with the literature on base-rate neglect. According to this, people, including physicians, neglect prevalence information (e.g., Elstein & Schwartz, 2000; Hamm, 1996; Hoffrage & Gigerenzer, 1998; Kahneman & Tversky, 1973; McNair & Feeney, 2015; Sirota, Juanchich, & Hagemayer, 2014; Sirota, Kostovičová, & Vallée-Tourangeau, 2015). For instance, only 5 out of 48 physicians integrated the disease base-rate information with the results of cancer screening, when asked about its positive predictive value (Hoffrage & Gigerenzer, 1998). However, we wish to argue that this is only a superficial contradiction, because of different contexts (screening vs. diagnosing), requiring different knowledge and engaging different cognitive processes. Neglecting the base-rate information seems to be limited to specific cognitive paradigms (e.g., textbook tasks) that require people to compute the probabilities explicitly and provide the statistical information from description, not from experience (Hertwig, Barron, Weber, & Erev, 2004; Sirota, Vallée-Tourangeau, Vallée-Tourangeau, & Juanchich, 2015). In fact, some authors would argue that people, including physicians, are sensitive to the base-rate information and integrate it successfully with other pieces of information once they have experienced them (Christensenszalanski & Beach, 1982; Gill, Sabin, & Schmid, 2005). In our study, we provided qualitative evidence of the updating that is based on experience.

The effect of an alternative explanation fits well with the hypothesis generation model (Thomas et al., 2014; Thomas et al., 2008). The immediate availability of an alternative explanation creates a stronger link between the relevant diagnostic hypothesis and the observed symptoms and enables it to reach the activation threshold for entering working memory. This would also fit well with the general cognitive tendency of people to focus too narrowly on the
hypothesis currently under consideration, while ignoring other relevant hypotheses. People tend to neglect relevant alternative explanations (Fernbach, Darlow, & Sloman, 2010), favor information supporting the focal hypothesis (Doherty, Chadwick, Garavan, Barr, & Mynatt, 1996), direct a searching strategy that confirms their initial hypothesis (Klayman & Ha, 1987) and construct one mental model of a diagnostic problem at a time and then focus on it (see singularity principle, Evans, 2006). Future research should test further the specific psychological mechanisms of hypothesis activation and maintenance in medical diagnosis, using more real-world study designs, e.g. where physicians are allowed to gather information as they see fit, rather than being provided with information all at once or in a predetermined sequence.

Physician experience increased the frequency of cancer provided as the main working diagnosis. This is, however, in direct contrast with the findings of a recent study (Kostopoulou et al., 2016), where family physicians diagnosed richer and more realistic cancer scenarios than the vignettes used in this study: more experienced physicians were significantly less likely to consider cancer as a possibility at the start of the clinical consultation, and to give it later as their main, working diagnosis. In this and the current study, nevertheless, the relationship between experience and diagnosis was weak, with odds ratios close to 1. These different findings exemplify the lack of consensus in the literature regarding the relationship between physician experience and diagnostic accuracy.

Our findings have theoretical implications. First, they provide some validation of the hypothesis generation model using a realistic task: physicians diagnosed and managed the hypothetical patients consistently with the probabilistic nature of this process, sensitive to changes in prevalence and to a more strongly-activated, competing diagnosis. These findings are also consistent with other theoretical models, such as the Extended Analytical-Heuristic
Processes Model (Evans, 2006). Prevalence of cancer likely plays some role in constructing the most plausible mental model of the symptoms; if the mental model of an alternative diagnosis is generated and meets the satisfaction principle threshold, then it is likely to be accepted; this, in turn, may eliminate the less likely mental model of cancer. However, these post-hoc explanations of our findings cannot corroborate Evans’ model, as it does not postulate – in contrast with the HyGene model – these specific predictions a priori.

Furthermore, we did not study an exhaustive list of factors postulated by the HyGene model that could affect the hypotheses generation phase and, in turn, cancer detection. Other cognitive factors, e.g., working memory capacity and the number of diagnostic alternatives considered, as well as situational factors that directly impinge on cognition, such as time constraints and order of presented data are of equal theoretical importance for model validation on practicing physicians, and should be included in future studies (e.g., Doherty et al., 1996; Dougherty, Gettys, & Thomas, 1997; Dougherty & Hunter, 2003a, 2003b; Thomas et al., 2014; Thomas et al., 2008). The two factors that we chose to test in this study are theoretically important for the model validation and could be adopted in future interventions to improve cancer detection, as suggested below.

Finally, some physicians generated cancer even when its prevalence was low and an alternative, more common explanation was available. This may indicate concern with less common but more serious disease. Although we did not measure physicians’ perceived probability of cancer, there is some evidence that the severity of the diagnostic outcome can influence hypothesis generation and inflate posterior probability judgments (Harris & Corner, 2011; Juanchich, Sirota, & Butler, 2012; Weber et al., 1993). This could be considered within the HyGene model – perhaps the traces encoding rare but costly diagnoses in episodic memory are
activated more easily and with minimal cue input.

Our findings have practical implications for medical education and training and the design of diagnostic support. In terms of education and training, if retrieval from episodic memory is crucial for early cancer diagnosis, then simulated experience rather than simple provision of statistical information may lead to better probability calibration (Hertwig et al., 2004). Appropriate calibration of probability may also be useful in avoiding over-diagnosis and over-referral for cancer. In terms of diagnostic support for hypotheses generation, offering a list of differential diagnoses at the beginning of the consultation, and no other help, was found to increase the final diagnostic accuracy of family physicians in two European countries (Kostopoulou, Lionis, et al., 2015; Kostopoulou, Rosen, et al., 2015). Specifically, family physicians in the UK and Greece diagnosed a series of simulated patients online. One group received early diagnostic support in the form of a list of diagnostic suggestions at the start of the consultation, after reading some limited information (a short patient description and the presenting problem) and before collecting further information to test their hypotheses. In both countries, the group who received early diagnostic support was significantly more accurate than the unaided control group (pooled OR 1.40, 95% CI [1.13-1.67]). Kostopoulou and colleagues subsequently designed a prototype for a Decision Support System, based on the principle of early diagnostic support, which they evaluated in a high-fidelity simulation (physicians consulting with actors-as-patients) and found significant improvement in diagnostic accuracy, without an associated increase in the number of investigations (Kostopoulou, Porat, Corrigan, S., & Delaney, in press). These studies provide evidence that early suggestions of diagnostic alternatives, before the physician engages in information gathering and hypothesis testing, can lead to improvements in diagnostic accuracy.
We acknowledge the tension between missing serious diseases, such as cancers, and over-diagnosis, leading to unnecessary investigations and sometimes treatment. Until there is more and better evidence to inform appropriate risk assessment, and until economic analyses are carried out to determine the most appropriate decision thresholds for referrals and investigations, such tensions will persist (Rubin et al., 2015). Although risk cancer tools are currently available, such as QCancer (http://www.qcancer.org) and The Risk Assessment Tool, which can calculate the patient’s cancer risk, employing them requires that cancer has already been considered as a possibility. In fact, the argument for early support, i.e., support to influence a decision being made rather than change a decision already made, has been articulated in relation to other medical decision tasks, such as prescribing (Hayward et al., 2013).

Our study has several methodological limitations: two of which deserve more attention. First, one could question the external validity of the vignettes used in this study. We believe that the vignettes represented the diagnostic and management decisions well, since our approach to the patient vignettes’ creation corresponded to the general recommendations for designing externally valid clinical judgment vignettes (e.g., Evans et al., 2015). For instance, the vignettes were constructed with the help of family physicians, after reviewing the relevant clinical literature, and included repeated patient presentations, as is common in patients with cancer. Furthermore, a recent multi-country study found that physicians’ willingness to investigate possible cancer symptoms in vignettes was associated with national survival rates (Rose et al., 2015). Nevertheless, future research should corroborate our findings in more realistic conditions, as suggested earlier. Second, one could question to what extent our findings could be generalized to other cancers. Indeed, we simulated only two common and well-known cancers. We would expect qualitatively similar processes to be at play in other cancers and – in light of current
findings – an amplified effect of missing early presentations of less common and less-known cancers.

In summary, family physicians diagnosed cancer more frequently when the prior probability of cancer in patients presenting with vague symptoms was higher, when they were not offered an alternative explanation of the symptoms, and when the patient presented repeatedly with worsening symptoms. Physicians’ working diagnoses guided their referral decisions. Our findings corroborate some of the assumptions of the hypothesis-generation model with practicing physicians, and, along with the other studies discussed, emphasize the importance of the initial diagnostic phase, when competing working diagnoses are generated. Future interventions aiming to improve cancer detection could target this phase, by offering possible differential diagnoses with severe consequences and/or with underestimated prior probabilities.
References


computer-simulated patients. British Journal of General Practice, 65(630), e49-54. doi: 10.3399/bjgp15X683161


### Table 1. Effect of prevalence and a prior alternative explanation on cancer diagnosis.

<table>
<thead>
<tr>
<th>Factors</th>
<th>Outcomes</th>
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<tr>
<td></td>
<td>Cancer as the main diagnosis</td>
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<tr>
<td></td>
<td>Unadjusted model</td>
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<tr>
<td></td>
<td>OR, 95% CI</td>
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<td></td>
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<tr>
<td>Alternative</td>
<td>1.70, [1.11, 2.59]</td>
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<td></td>
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<tr>
<td>Prevalence*</td>
<td>0.62, [0.35, 1.09]</td>
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<tr>
<td>Alt. Explanation</td>
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<td>Covariates</td>
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<td>Consultation</td>
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<td></td>
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<tr>
<td>Experience</td>
<td>1.03, [1.02, 1.05]</td>
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</table>

*Adjusted for consultation and experience

Note. OR = odds ratio; 95% CI = 95% confidence interval; z = z-test, p = statistical significance
Table 2: Effect of reporting cancer as the main working diagnosis on urgent cancer referral (two-week referral for cancer) and on any referral leading to cancer detection.

<table>
<thead>
<tr>
<th>Factors</th>
<th>Early cancer referral (vs. delay or no referral)</th>
<th>Cancer referral (vs no referral)</th>
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<tr>
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<td>Unadjusted model</td>
<td>Adjusted model</td>
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<tr>
<td></td>
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<td>OR, 95% CI</td>
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<td>Z = 15.71, p &lt; .001</td>
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<td>—</td>
<td>Z = 9.73, p &lt; .001</td>
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<td>Experience</td>
<td>0.99, [0.97, 1.01]</td>
<td>—</td>
</tr>
<tr>
<td>Z = -0.59, p = .555</td>
<td>—</td>
<td>Z = -0.46, p = .646</td>
</tr>
</tbody>
</table>

Note. OR = odds ratio; 95% CI = 95% confidence interval; Z = z-test, p = statistical significance
*Adjusted for consultation and experience
Figure Captions

*Figure 1.* Effects of the main factors on cancer given as the main diagnosis (in %) and on urgent referral for cancer (in %). Panel A: Effect of cancer type and patient consultation on generating cancer as the main diagnosis; Panel B: Effect of prevalence and alternative explanation on generating cancer as the main diagnosis; Panel C: Effect of cancer type and patient consultation on urgent referral for cancer; Panel D: Effect of prevalence and alternative explanation on urgent referral for cancer.

*Note.* The error bars represent 95% Agresti-Coull confidence intervals for a binomial proportion.