

# A half-second glimpse often lets radiologists identify breast cancer cases even when viewing the mammogram of the opposite breast

Karla K. Evans<sup>a,1</sup>, Tamara Miner Haygood<sup>b</sup>, Julie Cooper<sup>c</sup>, Anne-Marie Culpan<sup>d</sup>, and Jeremy M. Wolfe<sup>e,f,g</sup>

<sup>a</sup>Department of Psychology, University of York, Heslington, York YO10 5DD, United Kingdom; <sup>b</sup>Department of Diagnostic Radiology, University of Texas MD Anderson Cancer Center, Houston, TX 77030; <sup>c</sup>Department of Radiology, York Teaching Hospital, York YO31 8HE, United Kingdom; <sup>d</sup>Division of Biomedical Imaging, University of Leeds, Leeds LS2 9JT, United Kingdom; <sup>e</sup>Department of Ophthalmology, Harvard Medical School, Boston, MA 02115; <sup>f</sup>Department of Radiology, Harvard Medical School, Boston, MA 02115; and <sup>g</sup>Department of Surgery, Brigham and Women's Hospital, Boston, MA 02115

Edited by Michael I. Posner, University of Oregon, Eugene, OR, and approved July 19, 2016 (received for review April 18, 2016)

Humans are very adept at extracting the “gist” of a scene in a fraction of a second. We have found that radiologists can discriminate normal from abnormal mammograms at above-chance levels after a half-second viewing ( $d' \sim 1$ ) but are at chance in localizing the abnormality. This pattern of results suggests that they are detecting a global signal of abnormality. What are the stimulus properties that might support this ability? We investigated the nature of the gist signal in four experiments by asking radiologists to make detection and localization responses about briefly presented mammograms in which the spatial frequency, symmetry, and/or size of the images was manipulated. We show that the signal is stronger in the higher spatial frequencies. Performance does not depend on detection of breaks in the normal symmetry of left and right breasts. Moreover, above-chance classification is possible using images from the normal breast of a patient with overt signs of cancer only in the other breast. Some signal is present in the portions of the parenchyma (breast tissue) that do not contain a lesion or that are in the contralateral breast. This signal does not appear to be a simple assessment of breast density but rather the detection of the abnormal gist may be based on a widely distributed image statistic, learned by experts. The finding that a global signal, related to disease, can be detected in parenchyma that does not contain a lesion has implications for improving breast cancer detection.

gist processing | medical image perception | attention | mammography

Rapid extraction of scene “gist” (1–4) is a very useful aspect of routine visual perception that allows us to allocate our time and attention intelligently when confronted with new visual information (Can I find food here? Is there danger here?). The signals that we extract on our first glimpse of a scene are imperfect but not random. Experts often anecdotally report gist-like experiences with complex images in their domain of expertise. For instance, we have shown that radiologists can distinguish normal from abnormal mammograms at above-chance levels in as little as a quarter of a second, whereas nonexperts cannot (5). The gist of abnormality appears to be a global signal. Radiologists can detect it but cannot even crudely localize the abnormality under these conditions.

Detecting the gist of breast cancer might be more than a curiosity, if that signal could be used to improve performance in breast cancer screening. Screening mammography can reduce mortality through early diagnosis of disease (6). Breast cancer is the most prevalent cancer in women and is the second leading cause of cancer deaths in women (7). In North America, screening mammography has a false negative rate of 20 to 30% (8, 9) and a recall rate of about 10% (10). With a disease prevalence of about 0.3% (11), the vast majority of those recalled will not have cancer. Thus, there is significant room for improvement.

It has been argued for many years that an initial, global processing step is an important component in expert medical image

perception that might constrain or filter subsequent search (12–15), with the two most prominent models (16, 17) each placing great emphasis on experts' ability to process and evaluate information from large regions of an image (18). These models are broadly consistent with two-stage models of visual search (19, 20), developed in the basic vision literature that propose that there is a limited set of features that can be used to guide attention and subsequent serial stage that allows for “binding” of features to permit identification of objects. Global processing of scene gist is a component of a recent modification of this class of model (21). This formulation proposes there is a selective pathway that can be used to recognize one (or a very few) objects at a time. Access to this limited-capacity process is controlled by attention and the deployment of attention is guided by the basic features, mentioned above. There is also a nonselective pathway, capable of rapid extraction of “global image statistics” like the average orientation of a set of line segments or the average size of objects (22–24). Perhaps more interestingly, the distribution of basic features, the “spatial envelope” (25, 26), contains information that allows for semantic categorization of scenes (e.g., natural vs. urban) without the need to recognize specific objects in the scene.

It is important not to oversell the capabilities of the nonselective pathway. It is engaged in global processing and cannot reliably recognize specific objects. Moreover, the discriminations made on the basis of a first glimpse, while not random, are typically far from perfect. Returning to mammography, Evans et al. (5) found that, although experts could classify mammograms

## Significance

Discovering characteristics of a signal that indicates to medical experts the presence of cancer in a noninvasive screening technique in a blink of an eye has implications for improving cancer detection. Here we report two surprising facts about this signal. First, it is much stronger in the high spatial frequencies (fine detail) than in the low frequencies. Second, it is widely distributed, with signal being present well away from the actual visible locus of disease even in the breast contralateral to visible signs of disease. Although this signal is not, in itself, definitive, it has the potential to be used in automated aids to medical screening and incorporated into training protocols for medical experts, speeding up and improving cancer detection.

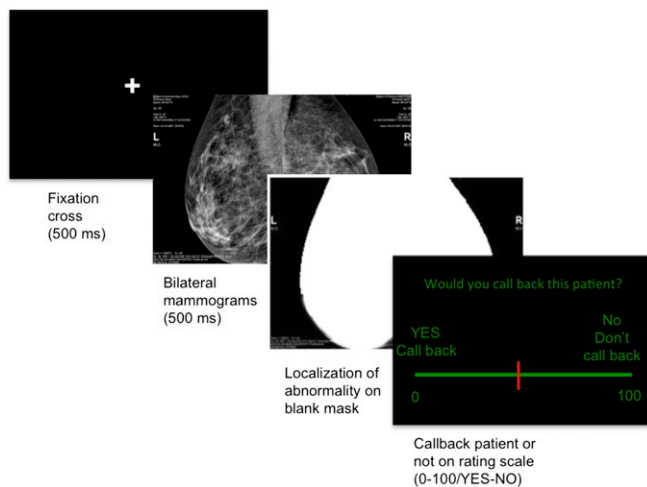
Author contributions: K.K.E. and J.M.W. designed research; K.K.E., T.M.H., and A.-M.C. performed research; K.K.E. and J.M.W. analyzed data; K.K.E., T.M.H., J.C., A.-M.C., and J.M.W. wrote the paper; and T.M.H. and J.C. provided necessary density rating and expertise in radiology.

The authors declare no conflict of interest.

This article is a PNAS Direct Submission.

<sup>1</sup>To whom correspondence should be addressed. Email: karla.evans@york.ac.uk.

This article contains supporting information online at [www.pnas.org/lookup/suppl/doi:10.1073/pnas.1606187113/-DCSupplemental](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1606187113/-DCSupplemental).



**Fig. 1.** Experimental procedure for experiments 1–4. Experiment 2 just showed a unilateral breast image, and experiment 4 used only a piece of the breast image.

as normal or abnormal at above-chance levels, they were at chance in their ability to localize abnormalities. Nevertheless, mammograms appear to contain a signal indicating abnormality. This profile of image statistics or global properties might guide attention or, at least, might alert the radiologist to the possible presence of an abnormality in a mammogram.

In this paper, we investigate the nature of this global signal in the hope that the signal could be better exploited by radiologists or used by designers of computer-aided detection (CAD) systems to improve breast cancer screening. Our results show that the signal is concentrated in the high spatial frequencies of the image. It is not based on symmetry between two breasts or density of the breasts. Finally, the signal is detectable in breast tissue away from the location of the actual abnormality, including in the contralateral breast. In each of four experiments, we presented experienced radiologists with unilateral or bilateral mammograms [craniocaudal (CC) or mediolateral oblique (MLO) views of both breasts] or sections of mammograms for 500 ms (allowing for, perhaps, two volitional fixations). The stimuli were followed by a mask (a white outline of the breasts). Observers rated each stimulus on a scale from 0 (certainly recall this patient) to 100 (certainly normal) (Fig. 1). If the stimulus was a full breast or pair of breast images, observers were asked to localize the abnormality on an outline of that breast image. We also obtained density ratings from other radiologists for the mammogram stimulus set used in the experiments (full methods are presented in *Methods*).

## Results

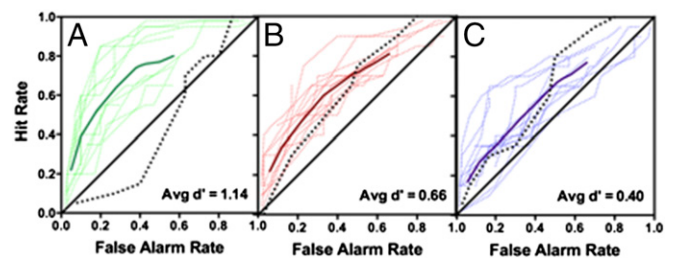
**Experiment 1.** Experiment 1 asked if the abnormality signal was based on a disruption in the usual bilateral symmetry of the breasts. Studies have noted that asymmetry can be a strong indicator for developing breast cancer (27, 28). Indeed, research has suggested that bilateral mammographic density asymmetry could be a significantly stronger risk factor for breast cancer development in the near-term than either woman's age or mean mammographic density (29).

We measured observers' ratings of abnormality to three types of images: (i) baseline, both breasts from the same woman; (ii) asymmetry 1, breast images from two different women (on positive/abnormal trials, one breast image was abnormal, whereas the other was a normal image from another woman); and (iii) asymmetry 2, breasts are from two different women (on positive trials, one breast image was abnormal with a lesion, whereas the other image came from the breast contralateral to

a lesion in another woman) (Fig. 2).  $d'$ , the signal detection measure of performance, is calculated by comparing ratings of the abnormal condition to the ratings of the otherwise equivalent normal condition. When both breasts came from the same woman, expert radiologists could reliably exceed chance performance [average  $d' = 1.14$ ,  $t(13) = 8.69$ ,  $P < 0.0001$ ]. When the two breast images came from two different women, radiologists could still perform the task [average  $d' = 0.66$ ,  $t(13) = 6.28$ ,  $P < 0.0001$ ], although their performance was significantly worse than when both breasts were from the same woman [planned comparison,  $t(13) = 7.03$ ,  $P = 0.018$ ]. When the abnormal case consists of one breast with an abnormality and the other breast was the breast contralateral to the lesion from a different woman, again, radiologists could do the task [average  $d' = 0.40$ ,  $t(13) = 3.02$ ,  $P < 0.00097$ ], but their performance was weaker than the performance in the condition where both breasts were from the same woman ( $P = 0.054$ ). Performance did not differ significantly between the two asymmetric conditions ( $P > 0.05$ ). We can conclude from these results that symmetry may be part of what allows an expert to distinguish a normal from abnormal case in a glance, but it is not required because there is above-chance performance in the artificial, asymmetric conditions.

Although participants could detect the presence of abnormality, they could not localize that abnormality when it was present (Fig. S1). Localization performance was not significantly different from chance. Localization was best for the baseline condition (21%), but still not above chance performance [ $t(13) = 1.38$ ,  $P = 0.196$ ]. In addition, as shown in Evans et al. (5), localization performance did not improve as the confidence rating increased.

Is the signal of abnormality simply breast density, with dense breasts rated as more abnormal? In the baseline condition,  $d'$  was significantly better [ $t(13) = 6.93$ ,  $P < 0.0001$ ] than the  $d'$  derived from density estimates made by other radiologists. In the asymmetry conditions, the observed  $d'$  was not significantly better than  $d'$  derived from density rating [ $t(13) = 1.84$ ,  $P = 0.089$ ;  $t(13) = 0.48$ ,  $P = 0.647$ ]. However, if observers were basing their abnormality ratings on an assessment of density, one would expect that the gist and density ratings would be correlated, which they are not ( $r = 0.02$ ). One might also expect a difference in density ratings between normal and abnormal images. However, there is no reliable difference in this image set. A one-way ANOVA on density rating revealed no effect of image type [ $F(4,115) = 1.55$ ,  $P = 0.19$ ], whereas a one-way ANOVA revealed a large effect of image type on abnormality rating [ $F(4,115) = 18.5$ ,  $P < 0.0001$ ]. Thus, although the magnitude of the effect in the asymmetrical cases is similar to

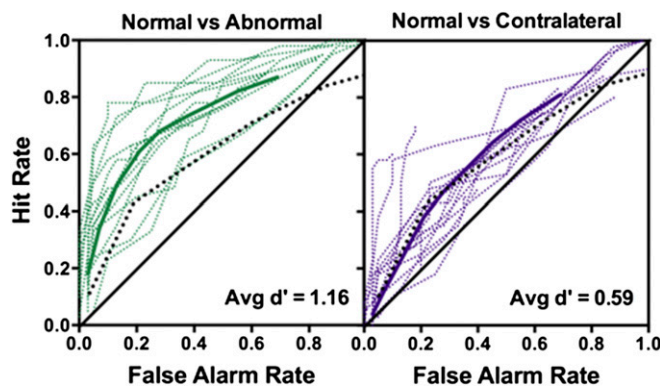


**Fig. 2.** Receiver operating characteristic (ROC) curves for the three conditions of experiment 1. Solid colored line, average ROC curve; light dotted lines, individual observers; dark dotted line, hypothetical ROC curve if judgments were based on density ratings. (A) Images of two breasts from the same woman. One breast abnormal on the positive trials.  $d' = 1.14$  compared with  $d' = 0.18$  derived from the density ratings. (B) Images from two different women, one image abnormal on the positive trials, and the other, always drawn from a negative case.  $d' = 0.66$  compared with  $d' = 0.47$  derived from the density ratings. (C) Images from two different women, one image abnormal on the positive trials, and the other image was the breast contralateral to a lesion in another woman.  $d' = 0.40$  compared with  $d' = 0.34$  derived from the density ratings.

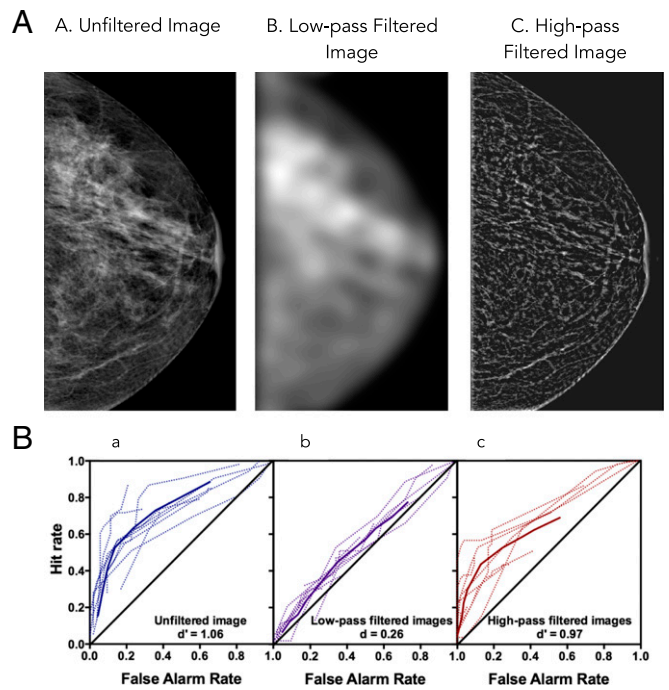
what could be obtained from a quick assessment of density, there is no evidence that density is the signal that was being used by our observers. Absence of evidence is not proof and it might be that a more statistically powerful experiment might show a relationship of perceived density and the gist of abnormality (e.g., an experiment with density and abnormality ratings made by the same observers). A different, perhaps simpler, way to test the symmetry question and to revisit the density question is to present radiologists with only brief presentation of a single breast image at one time, rather than with a paired viewing of the left and right breasts. That is the purpose of experiment 2.

**Experiment 2.** Participants rated the appearance of single breast images. In addition to determining whether observers can discriminate between normal and abnormal images in the absence of any possible symmetry signal, testing on single breast mammograms made it possible to assess whether the breast contralateral to an abnormal breast could be discriminated from breasts from negative cases. The left panel of Fig. 3 shows that observers were able to distinguish between images of single normal and abnormal breasts [ $d' = 1.16$ ;  $t(14) = 8.35$ ,  $P < 0.0001$ ]. What is more, as shown in the right panel of Fig. 3, their performance remained above chance when distinguishing normal from an image contralateral to the breast with a lesion [ $d' = 0.59$ ;  $t(14) = 8.35$ ,  $P < 0.0001$ ], although performance in that condition is significantly worse than performance with abnormal images [paired  $t(14) = 5.8$ ,  $P = 0.0004$ ]. As in experiment 1, the weaker performance, obtained with images contralateral to the lesion, was of a magnitude similar to what would be obtained if observers based their ratings on breast density. However, as in experiment 1, there is no evidence that the radiologists were using that density signal. As before, the relationship of density ratings to abnormality ratings was weak or nonexistent ( $r = 0.06$  for ratings and density across images and  $r = -0.02$  for the contralateral images alone). Further, there was no effect of the objective type of image (normal vs. abnormal) on density ratings [ $F(4,115) = 0.71$ ,  $P = 0.49$ ], but there was a large effect of image type on abnormality ratings [ $F(4,115) = 46.06$ ,  $P < 0.0001$ ]. As in experiment 1, the average localization performance of observers for images with the abnormality in a single breast was not significantly above chance level [ $t(14) = 0.91$ ,  $P = 0.378$ ].

**Experiment 3.** Any texture can be decomposed into a set of sinusoidal gratings of different spatial frequencies, amplitudes, orientations, and phases. Experiment 3 examined the spatial frequency composition of the signal of abnormality. Radiologists viewed normal and abnormal, bilateral mammograms in each of three counterbalanced conditions: unfiltered full images equivalent to



**Fig. 3.** ROC curves for single breast image data. Light dashed lines are individual observer data. The solid line shows the average data, and the dark dashed line shows the ROC curve that can be derived from the density data.



**Fig. 4.** (A) Example images used in experiment 3. (a) Unfiltered, (b) low-pass filtered, and (c) high-pass filtered views of a breast stimulus. (B) ROC curves for the three conditions of experiment 3. Solid colored lines, average ROC curves; dashed lines, individual observers. (a) Baseline, unfiltered/intact images. (b) Low-pass filtered images. (c) High-pass filtered images.

the baseline condition of experiment 1 and high-pass filtered images and low-pass filtered images shown as in Fig. 4A. There was a significant difference between conditions [ $F(2,16) = 52.35$ ,  $P < 0.0001$ ]. Specifically, the signal for interpreting mammography in 500 ms resides far more strongly in the high spatial frequencies, suggesting that the information is present in some aspect of the finer detail of the parenchymal texture (Fig. 4B). High-pass performance was reliably greater than chance [ $d' = 0.97$ ;  $t(8) = 8.05$ ,  $P < 0.0001$ ] and better than performance on low-pass images [low-pass  $d' = 0.26$ , paired  $t$  test,  $t(8) = 5.30$ ,  $P = 0.002$ ]. High-pass performance did not differ from performance with unfiltered images [ $d' = 0.97$  vs.  $1.06$ ,  $t(8) = 0.61$ ,  $P = 0.56$ ].

Again, the rated density of the images cannot explain radiologists' performance in any of the three conditions. The derived  $d'$  from the average density rating was  $d' = 0.09$  and that is significantly lower than the performance for unfiltered images [ $d' = 1.06$ ,  $t(8) = 8.81$ ,  $P < 0.0001$ ], high-pass images [ $d' = 0.97$ ,  $t(8) = 7.43$ ,  $P < 0.0001$ ], or low-pass images ( $d' = 0.26$ ,  $t(8) = 6.00$ ,  $P < 0.0003$ ). None of the correlations of image density and image abnormality rating were significant [all  $F(1,53) < 2.2$ , all  $P > 0.14$ ].

These findings are interesting for at least two reasons. First, if radiologists were simply using density as the signal, one might expect better performance from low spatial frequencies. Second, outside of radiology, the more typical finding in the appreciation of scene gist is that it is the low spatial frequency content that can be appreciated first in a brief flash; not the higher frequencies, although 500 ms would be long enough to appreciate both low and high frequencies in a typical scene gist experiment (30). Because localization performance remained poor across all conditions [best for high-pass filtered images but still not above chance,  $t(8) = 0.86$ ,  $P = 0.414$ ; Fig. S2], we conclude that it is not a specific detail of the lesion that is supporting the decision but rather, abnormality is judged based on some aspect of the overall texture that is best visualized in the higher spatial frequencies. Perhaps the signal is related to processes that create indications of disease like

spicules that might be enhanced in a high-pass view, but a larger dataset would be needed to test such a hypothesis.

**Experiment 4.** If the signal of abnormality is present throughout the parenchyma as would be predicted if that signal is truly a global signal, then it follows that a signal should be found in isolated regions of the breast that deliberately exclude the lesion. Alternatively, even though radiologists cannot explicitly localize abnormalities after a 500-ms flash, the signal might still arise exclusively from some small portion of the breast rather than being distributed widely. To test that hypothesis, in experiment 4, we presented 256- × 256-pixel patches of mammograms and asked radiologists to distinguish between normal and three types of potentially abnormal patches: patches containing the lesion, lesion-free patches from the abnormal breast, and lesion-free patches from the breast contralateral to the lesion. Observer's performance differed significantly between the three types of samples [ $F(2, 20) = 109.14, P < 0.0001$ ]. However, all three types of patches from abnormal cases could be distinguished from normal at above-chance levels. This significant difference can be seen by noting that virtually all of the individual observer data lies above the main diagonal, chance line in Fig. 5. Performance on sections with the lesions was significantly better than patches without the lesion from either the ipsilateral ( $P < 0.0001$ ) or contralateral breast ( $P < 0.0001$ ). Performance on ipsilateral and contralateral patches without a visible lesion did not differ ( $P = 0.473$ ). The density estimates, made by other radiologists for these small patches, produce area under the ROC curve (AUC) between 0.47 and 0.49, essentially at the 0.5 (chance) level. Apparently, there is no signal in the density ratings for these small patches of breast parenchyma. There was no difference between the average density ratings for the different types of sections [ $F(3,196) = 0.09, P = 0.97$ ], and the density ratings were not significantly correlated with the abnormality ratings (all  $r < 0.05$ , all  $P > 0.12$ ).

These results provide interesting insight into the signal supporting radiologists' performance in these tasks. Unsurprisingly, when the section includes the lesion, attention will be directed to the lesion, and performance is better than if the radiologist is looking at the entire breast with the lesion in an unknown location. Of more interest, there is some signal in sections of parenchyma ipsilateral and contralateral to the lesion. The signal is weak (Fig. 5, conditions B and C), corresponding to  $d'$  values of only 0.33–0.40 in the sections that did not include the lesion. However, note that the patches show only about one-eighth of a single breast. If we model the whole breast as consisting of eight independent samples with  $d' = 0.33$ –0.40, performance for a presentation of the whole breast would yield  $d'$  between 0.9 and 1.2. This modeled whole breast  $d'$  is comparable to or somewhat higher than the  $d'$  for whole breasts in experiments 1–3. If results

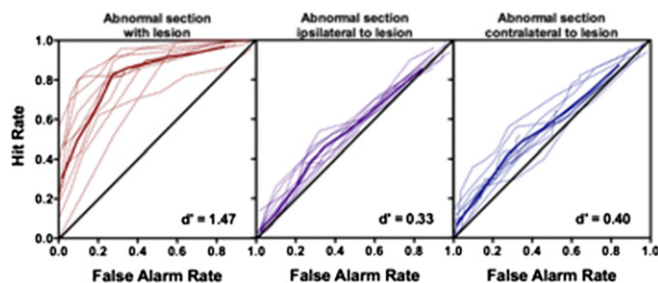


Fig. 5. ROC curves for the three conditions of experiment 4. Solid colored lines, average ROC curves; dashed lines, individual observers. (Left) Abnormal section contains lesion,  $d' = 1.47$  (99.9% CI, 1.20–2.12). (Middle) Abnormal section ipsilateral to lesion,  $d' = 0.33$  (99.9% CI, 0.17–0.49). (Right) Abnormal section contralateral to lesion,  $d' = 0.40$  (99.9% CI, 0.21–0.58). In all cases, the hit rate is derived from sections taken from a normal case.

from the whole breast are actually worse than would be predicted from small patches, that suggests that the signals combined across the whole breast are not entirely independent. In any case, the local signal is in principle, strong enough to support the results obtained with whole breasts, when combined across the whole breast.

## Discussion

Radiologists report anecdotally that some images seem to be “bad” when they first appear, before any specific pathology is localized. No one would suggest that diagnosis should be based on these first glimpses. However, there is now a body of research, including the work reported here, that indicates that this sense of the gist of a medical image can be based on a measurable signal (5, 12, 15). Our goal, in the present paper, has been to investigate the nature of the signal that allows expert observers to classify mammograms as normal or abnormal at above chance levels after a brief exposure. Experiments 1 and 2 undermined the hypothesis that observers were responding to a break in the normal rough symmetry between left and right breasts. In experiment 1, the symmetry was disrupted, and in experiment 2, observers only viewed a single breast image. In both cases, comparing normal and abnormal images, it remains possible to perform the classification task with a  $d'$  a bit better than 1.0. Although radiologists may use symmetry between two breasts as an important sign in normal mammography, it is not the signal that allows for classification of mammograms after a half-second of exposure.

Localization performance was consistently poor, suggesting that classification is based on a global signal, spread across the breast. The first interesting finding in this paper is the evidence in experiment 2 that this signal is present in the breast contralateral to the breast containing the abnormality. Experiment 4 found evidence for the signal in sections of parenchyma that did not contain an abnormality, regardless of whether they came from the ipsilateral or contralateral breast. Performance with these small sections is about what one would expect if the signal were being pooled across the entire image when the entire image is present. This finding may have clinical significance in the light of recent evidence that women with false-positive screening mammograms were at an increased risk of developing breast cancer compared with those with true negatives (31). Perhaps, even if localized signs of cancer were not unambiguously visible at the initial screening, radiologists still may have been influenced by the global signal of abnormality that we are studying here.

Experiment 3 provides another interesting finding: that the signal for abnormality is far stronger in a high-pass filtered mammogram than in a low-pass filtered image. Given prior results on recognition of briefly presented images (e.g., the global-local effect) (32, 33), one might have expected some sort of advantage for the coarser information in the low-pass image. Instead, we found the information about abnormality resides in the higher frequencies.

It is worth noting that the ability to detect abnormality at above-chance levels is a learned skill of expert radiologists. In previous work (5), we had nonexperts attempt the task. They performed at chance levels. It would be interesting to know if general radiologists who read fewer mammograms are able to detect this global signal of abnormality.

A distributed global signal of abnormality in breast cancer might be a useful component in a CAD system (34). The normal goal of a CAD system is to direct the radiologist's attention to specific, suspicious locations. Although these systems perform at a level comparable to that of an expert radiologist, they have not been hugely successful in clinical practice (35), in part because the positive predictive value of any given CAD mark is very low in a mammography screening situation where the prevalence of disease is low. As a result, radiologists tend to dismiss the correct CAD marks when they occur (36). It is possible that the signal that supports classification in the experiments reported here, could be used as an additional piece of information for a CAD

system. A CAD mark in the presence of a global abnormality signal might be a more suspicious mark than one in the absence of the signal. The presence of the signal in the breast contralateral to the abnormality also raises an interesting clinical possibility. It may be that the signal is present before the actual lesion appears. If so, it could be used as a warning sign, suggesting greater vigilance much as breast density is used as risk factor today (37). In thinking about any of these possibilities, it is critical to remember that radiologists' ability to detect abnormality in half a second is probabilistic. They perform above chance but far from perfect and far from their performance under normal conditions of reading mammograms. The gist signal might be useful, but, by itself, it is nowhere near definitive. In conclusion, there is a global signal that can be measured by asking radiologists to classify mammograms in a fraction of a second. That signal is probably the basis of the initial "holistic" impression of an image that is thought to guide radiologists when they view images in a normal, clinical setting (12, 38). If properly quantified, it could also be a component of automated aids to mammography.

## Methods

**Participants.** All study participants were attending radiologists specializing in breast imaging. Across the four studies, we tested 49 radiologists: experiment 1, 14 radiologists (11 female, 3 male; average age, 53 y), average 19 y in practice (range, 4–34 y) reading, on average, 7,650 cases in the last year (range, 6,000–10,000); experiment 2, 15 radiologists (12 female, 3 male; average age, 49 y), average 19 y in practice (range, 10–35 y), reading, on average, 7,280 cases in the last year (range, 3,000–15,000); experiment 3, 9 radiologists (5 female, 4 male; average age, 50 y), average 15 y in practice (range, 7–39 y), reading, on average, 7,100 cases in the last year (range, 4,000–10,000); and experiment 4, 11 radiologists (10 female, 1 male; average age, 52 y), average 20 y in practice (range, 4–34 y), reading, on average, 7,800 cases in the last year (range, 6,000–10,000). The radiologists who participated in experiments 1, 2, and 4 were recruited from five National Health Service (NHS) Hospital Trusts in Yorkshire and Cambria, UK. In experiment 3, radiologists were recruited from University of Texas MD Anderson Cancer Center (Houston). All of the participants had normal or corrected-to-normal vision and gave informed consent. The experiments had institutional review board approval from University of York, University of Texas MD Anderson Cancer Center, and the NHS Hospital Trusts.

**Stimuli and Materials.** The stimuli used in the four experiments were derived from 120 bilateral full-field digital mammograms. The starting resolution of the two mammograms side by side was  $1,980 \times 2,294$  pixels. These bilateral mammograms were then downsized to fit on a monitor with a resolution of  $1,920 \times 1,080$ . Mammograms were acquired from anonymized cases from Brigham and Women's Hospital (Boston). All of the cases included at least four images (left and right breast MLO views and CC views). Half of the cases showed cancerous abnormalities, whereas the rest were normal. Abnormal cases were either screen-detected cancers, histologically verified, or mammograms that had been done 1–2 y before a screen-detected cancer and that had been interpreted as negative but later retroactively determined by a study radiologist to have contained visible abnormalities. The abnormalities demonstrated on mammograms were "subtle" masses and architectural distortions. Lesion subtlety was determined by the study radiologists based on their experience. We did not include calcifications or more obvious cancers as it is of less interest to show that a stimulus like a bright white spot can be detected in less than a second. The average size of the lesions in the test set mammograms was 18 mm (range, 10–48 mm).

Experiments 1 and 3 used all of the 120 bilateral mammograms. For experiment 2, these original images were Fourier transformed, and two types of filtered images were computed. A low-pass image was created by removing all of the information above two cycles per degree (at a 57-cm viewing distance), leaving only the low spatial frequencies of the original image. A high-pass image was created by removing information that was below six cycles per degree, leaving only the high spatial frequency information of the original images. Application of Fourier transform of images resulted in three sets of images: original intact images, the same images but with only low spatial frequency information present, and images with only the high spatial frequency information present.

In experiment 2, we used 120 unilateral breasts, taken from the bilateral full-field digital mammograms used in experiment 1. A third of the single

mammograms had a confirmed yet subtle abnormality (e.g., mass or architectural distortion), another third was taken from completely normal cases, and the last third was mammograms of breasts that contained no abnormality but that were the breast contralateral to a breast containing an abnormality.

The stimuli used in experiment 4 consisted of 200 sections taken from the original full-field digital mammograms (including both CC and MLO views). Mammogram sections were cropped to  $256 \times 256$  pixels using Photoshop CS6 (Adobe). A quarter of the sections included a lesion, centered in the patch. There were three types of no lesion sections: (i) section taken from the image of an abnormal breast but not containing the lesion, (ii) section taken from the breast contralateral to a breast containing a lesion, and (iii) section taken from a completely normal case.

Two of the authors (T.M.H. and J.C.), who are practicing radiologists, provided density ratings for each left and right mammographic image for all of the images in the stimulus set on a four-point scale (1, fatty; 2, scattered fibroglandular; 3, heterogeneously dense; 4, extremely dense). The density ratings of the two radiologists were significantly correlated for both breasts (left breast:  $r = 0.56$ ,  $P < 0.00001$ ; right breast:  $r = 0.43$ ,  $P < 0.00001$ ). Rated density of abnormal cases was slightly higher than for normal [2.80 vs. 2.65, but not significantly, one-tailed  $t$  test  $t(188) = 1.64$ ,  $P = 0.101$ ]. If classification of normal vs. abnormal was based on the average density ratings given by the two radiologists, the predicted  $d'$  would be 0.26. The density ratings of the two radiologists for the single breast subset of stimuli used in experiment 2 were also significantly correlated ( $r = 0.36$ ,  $P < 0.0001$ ). The density raters also gave a density rating for the four types of section we used in experiment 4. A one-way independent ANOVA on the average density rating revealed no significant main effect of type of section [ $F(1,199) = 0.86$ ,  $P = 0.968$ ]. Thus, there was no significant difference in the density rating of the four types of small section.

Experiments 1, 2, and 4 were conducted on a Macintosh MacBook Pro using MATLAB R2012b. All observers viewed the experiment on a 27.5-in, liquid-crystal color screen with a  $1,920 \times 1,080$  resolution, a usable intensity range of 2–260 candelas per square meter, a contrast ratio of 188:1, and a refresh rate of 144 Hz at a viewing distance of 57 cm. Experiment 3 was conducted on a Dell Precision M6500 laptop using MATLAB R2012b. The experiment was displayed on a 17-in. screen at a viewing distance of 57 cm. The display monitor had a resolution of  $1,920 \times 1,200$  (Dell) and a refresh rate of 85 Hz.

**Procedure.** Across the study, all four experiments used the same experimental paradigm of brief stimuli presentation. All observers in each experiment viewed the same images, with the order randomized across trials. After three to six practice trials, depending on the experiment, each trial consisted of the following sequence of events. First, a fixation cross appeared in the center of the screen for 500 ms. The fixation cross display was followed by the brief 500-ms presentation of a pair of mammograms (experiments 1 and 3), side by side; a single mammogram (experiment 2); or a single section (experiment 4). After the brief presentation, observers saw a white outline of the previously presented breasts (experiments 1–3) or a white noise mask for another 500 ms (experiment 4). In experiments 1–3, even if they did not think the case was abnormal, radiologists were asked to indicate the most likely location of an abnormality with a mouse click on the display screen. Following this, observers were asked to provide a 0–100 rating (where 0 stands for clearly abnormal) scale how likely it was that there was an abnormality. Feedback was provided only for the initial practice trials. All of the observers were alone when performing the experiment.

In experiment 1, participants completed 120 trials across five possible trial types: (i) the two breasts were from the same woman, one breast with an abnormality, one normal (20 trials); (ii) the two breasts were from two different women, one breast with an abnormality, the other normal and from a completely normal case (20 trials); (iii) the two breasts were from different women, one with an abnormality, the other normal but from an abnormal case (20 trials); (iv) the two breasts were from the same woman: both breasts normal (30 trials); and (v) finally, the two breasts were from different women, both breasts and both cases completely normal (30 trials). Thus, overall, half of the cases were normal and half were abnormal. These five types of presentation were used to create the three comparisons described in the results. A comparison of conditions i and iv replicates the previous work on detection of the gist of abnormality (baseline). Comparisons of conditions ii and iii with condition v (asymmetry 1 and 2) test for the presence of a nonselective, gist signal when a symmetry cue cannot be used.

In experiment 2, participants completed 120 trials evenly divided between images of three types of breast: normal, abnormal, and contralateral (being the normal breast contralateral to an abnormal breast).

In experiment 3, participants completed three blocks of 120 trials, for a total of 360 experimental trials in which they viewed CC or MLO views of

mammograms. In each block, the observers saw only one set of images: the original intact image set, the low spatial frequency image set, or the high spatial frequency image set. The viewing order of the blocks was counter-balanced across observers.

In experiment 4, observers completed two blocks of 100 experimental trials each in which they viewed sections of mammograms evenly divided between the four types described above.

#### Data Analysis.

**Assessing detection performance.** The observers in all four experiments gave confidence ratings on a scale from 0 (clearly abnormal) to 100 (clearly normal). For a given rating threshold, scores above that rating can be considered true negatives, if the stimulus is normal and miss or false-negative errors if the stimulus is abnormal. Scores below the level are deemed hits or true positives, if the case is abnormal and false alarm or false-positive errors if the case is normal. Categorizing responses in this manner for a range of values sweeps out a ROC curve. Thus, signal detection measures of  $d'$ , criterion, and AUC can be derived. For purposes of calculating  $d'$ , we used a rating threshold of 50.

The analysis of experiment 1 is somewhat complicated because there are three types of abnormal cases (trial types *i*, *ii*, and *iii*) and two types of normal cases (trial types *iv* and *v*). For each of the three critical comparisons of normal and abnormal, the hit rate is derived from one of the three abnormal conditions, and the false alarm rate is derived from one of the two normal conditions. When the abnormal cases are those in which left and right images were from the same woman (trial type *i*), the false alarm rate is derived from the normal cases in which left and right images are also from one woman (trial type *iv*). When the abnormal cases are those in which the left and right images are taken from the mammograms of different women (trial types *ii* and *iii*), the normal cases are, likewise, taken from cases in which the left and right images come from different women (trial type *v*).

For experiments 2 and 4, the average  $d'$  and ROC curves were created by taking the false alarm rates from the single breast or sections taken from

normal breasts and pairing them with the hit rate from each of the two potentially abnormal conditions in experiment 2 or three potentially abnormal conditions in experiment 4.

**Calculating density  $d'$ .** A value of  $d'$  can also be calculated from the average density ratings using the same method as described above for the abnormality rating. Breast density was rated on a four-point scale from 1 = fatty to 4 = extremely dense. For normal images, using a threshold of 2.5, if the density rating was above threshold, that rating would be categorized as a false positive. If it was below, it was deemed to be a true negative. For abnormal mammograms, if the density rating was above the 2.5 cutoff, then it was categorized as a hit; if below, it was a miss. We used values above threshold as the analog of target present (abnormal) response because previous research has found that increased density is associated with higher likelihood of cancer (29).

**Assessing localization performance.** To assess localization performance the observers were asked to click on an outline mask of the breast to indicate where they thought an abnormality was most likely to have been located. Localization performance was measured by determining what percentage of observers' clicks fell into the predetermined regions of interest (ROIs) centered on abnormalities. We then calculated the percentage of correctly localized abnormalities in respect to the overall number of abnormalities. Chance levels for localization performance were determined as average percentage of the breast encompassed by the ROI (abnormal region). Different abnormal cases would have larger or smaller ROIs. Averaged across cases with lesions, the ROI area was 18% in experiment 1; experiment 2 = 6%; experiment 3 = 16%. These values then represent the chance of hitting an ROI by placing a random mark on the breast outline.

**ACKNOWLEDGMENTS.** This work was supported in part by a grant from the John S. Dunn, Sr., Distinguished Chair in Diagnostic Radiology of University of Texas MD Anderson Cancer Center. J.M.W. was supported by NIH Grant EY017001.

1. Biederman I, Rabinowitz JC, Glass AL, Stacy EW, Jr (1974) On the information extracted from a glance at a scene. *J Exp Psychol* 103(3):597-600.
2. Potter MC, Faulconer BA (1975) Time to understand pictures and words. *Nature* 253(5491):437-438.
3. Li FF, VanRullen R, Koch C, Perona P (2002) Rapid natural scene categorization in the near absence of attention. *Proc Natl Acad Sci USA* 99(14):9596-9601.
4. Kirchner H, Thorpe SJ (2006) Ultra-rapid object detection with saccadic eye movements: Visual processing speed revisited. *Vision Res* 46(11):1762-1776.
5. Evans KK, Georgian-Smith D, Tambouret R, Birdwell RL, Wolfe JM (2013) The gist of the abnormal: Above-chance medical decision making in the blink of an eye. *Psychon Bull Rev* 20(6):1170-1175.
6. Smith RA, Cokkinides V, Eyre HJ; American Cancer Society (2004) American Cancer Society guidelines for the early detection of cancer, 2004. *CA Cancer J Clin* 54(1):41-52.
7. Siegel R, Naishadham D, Jemal A (2013) Cancer statistics, 2013. *CA Cancer J Clin* 63(1):11-30.
8. Bird RE, Wallace TW, Yankaskas BC (1992) Analysis of cancers missed at screening mammography. *Radiology* 184(3):613-617.
9. Majid AS, de Paredes ES, Doherty RD, Sharma NR, Salvador X (2003) Missed breast carcinoma: Pitfalls and pearls. *Radiographics* 23(4):881-895.
10. Gur D, et al. (2004) Changes in breast cancer detection and mammography recall rates after the introduction of a computer-aided detection system. *J Natl Cancer Inst* 96(3):185-190.
11. Lee CS, Bhargavan-Chatfield M, Burnside ES, Nagy P, Sickles EA (2016) The National Mammography Database: Preliminary Data. *AJR Am J Roentgenol* 206(4):883-890.
12. Kundel HL, Nodine CF (1975) Interpreting chest radiographs without visual search. *Radiology* 116(3):527-532.
13. Carmody DP, Nodine CF, Kundel HL (1981) Finding lung nodules with and without comparative visual scanning. *Percept Psychophys* 29(6):594-598.
14. Oestmann JW, et al. (1988) Lung lesions: Correlation between viewing time and detection. *Radiology* 166(2):451-453.
15. Mugglestone MD, Gale AG, Cowley HC, Wilson ARM (1995) Diagnostic performance on briefly presented mammographic images. *Medical Imaging 1995*, ed Kundel HL (International Society for Optics and Photonics, San Diego), pp 106-115.
16. Nodine CF, Kundel HL (1987) The cognitive side of visual search in radiology. *Eye Movements: From Psychology to Cognition*, eds O'Regan JK, Levy-Shoen A (Elsevier Science, Amsterdam), pp 572-658.
17. Swenson RG (1980) A two-stage detection model applied to skilled visual search by radiologists. *Percept Psychophys* 27(1):11-16.
18. Reingold EM, Sheridan H (2011) Eye movements and visual expertise in chess and medicine. *Oxford Handbook on Eye Movements*, eds Liversedge SP, Gilchrist ID, Everling S (Oxford Univ Press, Oxford), pp 528-550.
19. Treisman A (2006) How the deployment of attention determines what we see. *Vision Res* 46(8):411-443.
20. Wolfe JM (2007) Guided Search 4.0: Current progress with a model of visual search. *Integrated Models of Cognitive Systems*, ed Gray WD (Oxford Univ Press, New York), pp 99-119.
21. Wolfe JM, Võ MLH, Evans KK, Greene MR (2011) Visual search in scenes involves selective and nonselective pathways. *Trends Cogn Sci* 15(2):77-84.
22. Ariely D (2001) Seeing sets: Representation by statistical properties. *Psychol Sci* 12(2):157-162.
23. Chong SC, Treisman A (2003) Representation of statistical properties. *Vision Res* 43(4):393-404.
24. Robitaille N, Harris IM (2011) When more is less: Extraction of summary statistics benefits from larger sets. *J Vis* 11(12):18-18.
25. Oliva A (2005) Gist of the scene. *Neurobiol Attention* 696(64):251-258.
26. Oliva A, Torralba A (2006) Building the gist of a scene: The role of global image features in recognition. *Prog Brain Res* 155:23-36.
27. Scutt D, Lancaster GA, Manning JT (2006) Breast asymmetry and predisposition to breast cancer. *Breast Cancer Res* 8(2):R14.
28. Sun W, et al. (2014) Prediction of near-term risk of developing breast cancer using computerized features from bilateral mammograms. *Comput Med Imaging Graph* 38(5):348-357.
29. Zheng B, et al. (2012) Bilateral mammographic density asymmetry and breast cancer risk: A preliminary assessment. *Eur J Radiol* 81(11):3222-3228.
30. Schyns PG, Oliva A (1994) From blobs to boundary edges: Evidence for time- and spatial-scale-dependent scene recognition. *Psychol Sci* 5(4):195-200.
31. Henderson LM, Hubbard RA, Sprague BL, Zhu W, Kerlikowske K (2015) Increased risk of developing breast cancer after a false-positive screening mammogram. *Cancer Epidemiol Biomarkers Prev* 24(12):1882-1889.
32. Navon D (1977) Forest before trees: The precedence of global features in visual perception. *Cognit Psychol* 9(3):353-383.
33. Kimchi R (1992) Primacy of wholistic processing and global/local paradigm: a critical review. *Psychol Bull* 112(1):24-38.
34. Gierach GL, et al. (2014) Relationships between computer-extracted mammographic texture pattern features and BRCA1/2 mutation status: A cross-sectional study. *Breast Cancer Res* 16(4):424.
35. Cole EB, et al. (2014) Impact of computer-aided detection systems on radiologist accuracy with digital mammography. *AJR Am J Roentgenol* 203(4):909-916.
36. Nishikawa RM, Giger ML, Jiang Y, Metz CE (2012) Re: Effectiveness of computer-aided detection in community mammography practice. *J Natl Cancer Inst* 104(1):77-77, author reply 78-79.
37. McCormack VA, dos Santos Silva I (2006) Breast density and parenchymal patterns as markers of breast cancer risk: A meta-analysis. *Cancer Epidemiol Biomarkers Prev* 15(6):1159-1169.
38. Kundel HL, Nodine CF, Conant EF, Weinstein SP (2007) Holistic component of image perception in mammogram interpretation: Gaze-tracking study. *Radiology* 242(2):396-402.