Abstract

In this work we investigate whether the innate visual recognition and learning capabilities of untrained humans can be used in conducting reliable microscopic analysis of biomedical samples toward diagnosis. For this purpose, we designed entertaining digital games that are interfaced with artificial learning and processing back-ends to demonstrate that in the case of binary medical diagnostics decisions (e.g., infected vs. uninfected), with the use of crowd-sourced games it is possible to approach the accuracy of medical experts in making such diagnoses. Specifically, using non-expert gamers we report diagnosis of malaria infected red blood cells with an accuracy that is within 1.25% of the diagnostics decisions made by a trained medical professional.

Introduction

Crowd-sourcing is an emerging concept that has attracted significant attention in recent years as a strategy for solving computationally expensive and difficult problems [1–6]. In this computing paradigm, pieces of difficult computational problems are distributed to a large number of individuals. Each participant completes one piece of the computational puzzle, sending the results back to a central system where they are all combined together to formulate the overall solution to the original problem. In this context, crowd-sourcing is often used as a solution to various pattern-recognition and analysis tasks that may take computers long times to solve. One of the underlying assumptions of such an approach is that humans are better than machines at certain computational and pattern recognition tasks.

There has been much work in the general field of ‘gaming’ as a method for crowd-sourcing of computational tasks [1,7–13]. Digital games have been used as effective means to engage an individual’s attention to computational tasks of interest. If a pattern-recognition task can be embedded as part of an engaging game, then a gamer may help in solving this task together with other gamers. Recently a number of gaming platforms have been created to tackle problems in e.g., biology and medical sciences, allowing non-experts to take part in solving such problems. FoldIt [7–8], as an example, is a game in which players attempt to digitally simulate folding of various proteins, helping researchers to achieve better predictions about protein structures. EteRNA [9] is another game, which likewise makes use of crowds to get a better understanding of RNA folding.

In this work, we take a similar strategy and demonstrate a platform to use digital gaming and machine learning to crowd-source the analysis of optical microscopy images of biomedical specimens through engaging the interest of human game players (i.e., gamers). The primary goal of this methodology is to accurately diagnose medical conditions, approaching the overall accuracy of medical experts, while only using non-expert gamers (see Figure 1). The same method can also function as a telemedicine platform, where trained medical experts could be made part of our gamer crowd through various incentives.

In general there is much detail and subtlety associated with medical images, and therefore accurate analysis and interpretation of such images often become tedious and time consuming, even for highly trained professionals. Crowd-sourcing of microscopic analysis and related diagnosis through gaming is rather timely in several ways. First, with rapid advances in mobile telecommunication and internet technologies such as mobile-phones, tablet PCs, etc., we have hundreds of millions of active users and potential gamers in the cloud that are all connected to a global network. In addition to this massive crowd volume, over the last few years, there has been a significant effort to create cost-effective, compact and lightweight microscope designs such that even mobile-phones could be converted into microscopic analysis tools [14–21]. Similar to the development of the PC, this is a very important development since it could enable wide-spread use of optical microscopy globally, with several orders of magnitude increase in the number of microscope users over the next decade. As will be further discussed in this manuscript, these recent advances make it feasible to create a self-learning platform that leverages crowd-sourcing and gaming concepts to conduct...
accurate and sensitive microscopic analysis of biomedical specimens.

We believe that this crowd-sourcing and gaming based micro-analysis platform could in particular be significant for telemedicine applications such that diagnostics decisions can be remotely made without the need for a local medical expert (e.g., a pathologist), especially impacting the medical infrastructure in resource-poor countries. We hypothesise that the gaming community can develop better recognition skills over time for specific medical conditions through a scoring system built in the games to identify such abilities of individuals (some of whom may also be health-care professionals that e.g., are even paid for each image that is diagnosed as part of the game). In addition to providing accurate remote diagnosis, the datasets of biomedical images characterized through this distributed platform can potentially be used as training images for automated machine learning based algorithms that over time self-learn to make reliable diagnosis (Figure 1). As such, through this crowd-sourcing and gaming platform we can create a self-learning integrated network of microscopes and imagers toward intelligent automated biomedical micro-analysis and diagnosis. Once scaled up, this smart network may have a significant impact on e.g., medical, environmental, and biological sciences, among others, through various innovative uses of this network and its constantly expanding database. For instance, by creating large data libraries of various specimens (e.g., microbial communities, parasites, pathology slides, blood/sputum/pap smears etc.) we may be able to dynamically track both temporal and spatial evolution of different pathogens, diseases, or infectious outbreaks and be able to better investigate and identify the cause-effect relationships of these spatiotemporal patterns (see Figure 1).

To demonstrate the proof-of-concept of the above outlined framework, we chose malaria as the medical condition to be diagnosed, and developed a crowd-sourcing and distributed gaming platform that allows individuals from anywhere in the world to assist in identifying malaria infected red blood cells (RBCs) imaged under light microscopes. In addition, we developed an automated algorithm for diagnosing the same images using computer vision, and created a novel hybrid platform for combining human and machine resources toward efficient, accurate and remote diagnosis of malaria.

For this initial demonstration, we chose malaria since it is still a major health problem in many tropical and sub-tropical climates, including much of sub-Saharan Africa (see Figure 1). It is the cause of ~20% of all childhood deaths in this region, and almost 40% of all hospitalizations in whole of Africa. For diagnosis of malaria, conventional light microscopy remains the gold standard method. A pathologist must typically check between 100 and 300 different field-of-views (FOVs) of a thin blood smear (corresponding to inspection of at least 1,000 individual RBCs) using a light microscope with 100X objective lens before being able to reliably call a thin smear sample negative (i.e. not infected). This, however, is a very time-consuming task and a significant challenge given the large number of cases observed in these resource-poor settings (see Figure 1). Furthermore, approximately 60% of the cases reported in sub-Saharan Africa are actually false-positives,[22] leading to unnecessary treatments and hospitalizations.

For the same purpose of malaria diagnosis in resource poor conditions, rapid diagnostic tests (RDTs) are also being developed to create an alternative to optical microscopy. However, RDTs
still have various shortcomings such as: \(i\) being relatively expensive compared to microscopy, costing \(\sim 0.3 - 2.0\) USD per test; \(ii\) existence of insufficient information on their quality and the lack of ability to test their performance in the field, [i.e., the lack of quality controls]; \(iii\) poor heat and humidity stability; and \(iv\) mistrust of RDTs by health-care workers and community members [which is technically coupled to issues \(i\) and \(iv\) mentioned above] [22]. Therefore, microscopic imaging of blood smears still remains as the gold standard method toward diagnosis of malaria.

There have been prior attempts based on machine vision algorithms to automate the process of malaria diagnosis in Giemsa-stained thin blood smears using optical microscopy images with promising performance results [23–27]. However, there are a number of factors that can negatively affect the performance of such algorithms, including variations in blood smear preparation and cell density on the slide, as well as variations in illumination, digital recording conditions, optical aberrations and the lens quality. As a result, these methodologies have not yet been able to find their ways into mainstream malaria diagnostics tools to start replacing manual inspection of blood smears.

The human visual system, however, does not suffer from the above mentioned limitations, and can correctly recognise patterns of parasitic infection even under severe variations in sample preparation, density and imaging/recording conditions. As such, we believe that scaling up accurate, automated, and remote diagnosis of malaria through a crowd-sourcing and gaming platform may achieve significant impact in the developing world through: \(i\) Elimination of the overuse and misuse of anti-malarial drugs, which is quite important for avoiding long-term drug resistance issues; \(ii\) Improved management of non-malaria fevers by allowing malaria to be ruled out, so that patients can receive more appropriate treatment; \(iii\) Much better use of existing funds and reduction in drug stock-outs; and \(iv\) Reduced risks due to long-term side-effects of anti-malarial drugs on patients who do not actually need treatment. Finally, we should also emphasize that the same crowd-sourcing and gaming based micro-analysis and medical diagnosis platform could further scale up for a variety of other biomedical and environmental applications where microscopic images need to be examined by experts.

**Methods**

**Game Design**

We developed a digital gaming platform through which we allow an unlimited number of gamers from any location in the world to access and diagnose images of human RBCs that are potentially infected with *P. falciparum*. The game (see Figure 2) was implemented to be run both on PCs (using Adobe Flash running on any internet browser such as Internet Explorer, Mozilla Firefox etc.) and on Android-based mobile devices, including mobile-phones as well as tablet PCs. We preferred an open-source operating system for mobile devices such that other game developers could easily contribute to the same crowd-sourcing platform in the future.

Before starting to play the game, each gamer was given a brief online tutorial explaining the rules of the game and how malaria infected RBCs typically look with some example images. After this, each gamer played a training game where s/he was required to successfully complete in order to continue playing the rest of the game. This test game consisted of 261 unique RBC images, where 20 of them were infected. The gamers were required to achieve >99% accuracy in this training game, and in the case of failure, they were asked to re-play it until they achieved 99%. This way all the gamers became familiar with the rules of the game and were briefly trained on the diagnostics task. Note that this training game was required only once at the time when the gamers registered on our platform. Upon registration, a unique user ID was assigned to each gamer and her/his individual diagnostics performance was tracked. Furthermore, this training game provided direct feedback to the players on their performance and their mistakes through a scoring mechanism. Since the labels (i.e., infected cell vs. healthy cell) of all the images were known a priori for the purposes of this training game, the player’s score was updated throughout the game (i.e., positive score for correct diagnosis, and negative score for incorrect diagnosis). It is important to note that there exists a large body of work on educational games [28–29]. However, given that our focus was not to educate the players, and in fact it was to demonstrate the quality of diagnostic results that can be achieved through untrained (non-expert) individuals, this initial test/training game was designed in a simple repetitive fashion.

**Figure 3. Overview of the gaming analysis framework.** The images are treated as a sequence of binary values that are broadcast by the server. The gamers are effectively noisy repeaters that in the most ideal case output the correct symbol for the inputs that they receive. Each repeater transmits its own noisy version of the same input symbol to a decoder. The decoder combines all the received repeater outputs and decodes a final output \(z_i\) which ideally will be the correct label/diagnosis for the input images. The repeaters can be modelled as Binary Communication Channels (top-left). \(p_j\) corresponds to the probability of receiving symbol \(j\) when in fact symbol \(i\) was transmitted.

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As the gamer goes through the game, s/he is presented with multiple frames of RBC images. The gamer has the option of using a ‘syringe’ tool to ‘kill’ the infected cells one by one, or use a ‘collect-all’ tool to designate all the remaining cells in the current frame as ‘healthy’, which significantly speeds up the cell diagnosis process since most of the RBCs are healthy. Within each frame, there are a certain number of cells whose labels (infected or healthy) are known to the game, but unknown to the gamers. These control cell images allow us to dynamically estimate the performance of the gamers (in terms of correct and incorrect diagnosis) as they go through each frame and also help us assign a score for every frame that they pass through (Note that this is different in the training game where all the images are effectively control images). Once a frame is completed, a score is assigned based on the performance of the gamer only on the control images. These control images (roughly 20% of all the images) along with the scoring system allow the game to provide some feedback to the gamer on their performance such that as the gamers continue to play, they can improve their diagnostics performance. The images and their order of appearance were identical among different gamers, thus allowing us to make a fair comparison among their relative performances.

### Image Database

To build a malaria infected RBC database, we used thin blood smear slides that contained mono-layers of cultured human RBCs which were infected by *Plasmodium falciparum* (*P. falciparum*) forming the source for our image dataset (refer to Reference [17] for further details). These malaria slides were then scanned with a bright-field optical microscope using a 100X oil-immersion objective.
objective lens (numerical aperture: 1.25). At each FOV, the captured RBC images were passed on to an infectious disease expert for identification of *P. falciparum* signatures and digital labeling of each RBC image (positive vs. negative). This process generated a dataset of 7116 unique RBC images, with 1603 of them infected by the malaria parasite. To form the set of images to be used in our games, each individual RBC image was cropped and resized to fixed dimensions of 50×50 pixels. To further increase the total number of images and their diversity (in terms of sample preparation, density and imaging conditions), we also used a set of images provided by the Center for Disease Control (CDC), yielding an addition of 118 infected and 595 uninfected RBC images. With this, we had a total of 7829 characterized human RBC images, with 1721 of them infected with *P. falciparum*, forming our ground truth database for evaluating our crowd-sourcing, gaming, and machine-vision based diagnostics platform. Please refer to Text S1 for further details and Figure S4 for sample images. No IRB approval was required since the digital red blood cell images that we used in our work were not linked to any patient data or diagnosis and were digitally created and shared for microscopic training purposes.

**Diagnostic Analysis**

When analysing the game results, we have access to the individual performance parameters and diagnoses (for both the control images and the unknown test images). We fuse the results from all gamers that have completed a particular game and generate a more accurate set of diagnoses for the test RBC images. Given that each RBC image either corresponds to a healthy cell or an infected cell, we can use binary labels to identify them: 0 for healthy and 1 for infected. Recasting our setup as a communications system, our server will act as a broadcaster of a binary sequence and each gamer will act as a noisy Binary Channel [30], retransmitting the symbols back along with some errors. Therefore, we model the framework of our games as a noisy communication network consisting of a broadcast unit, multiple repeaters, and a receiver/decoder unit for the final diagnosis (see Figure 3). In the ideal scenario, the repeaters (i.e. the gamers) would simply receive a set of incoming symbols (images to be diagnosed) from the broadcast unit (through various light microscopes located in e.g., point-of-care offices or malaria clinics), and transmit them to the receiver/decoder block, which in turn computes the optimal “correct” label for each individual unknown RBC image using a Maximum a Posteriori Probability (MAP) approach. In Text S1 (Section II) we provide a theoretical description of how this performance analysis is done and is used to diagnose unknown RBCs based on gamers’ responses.

**Results and Discussion**

To test the viability of our crowd-sourced gaming-based malaria diagnosis platform, different experiments were run with 31 unique participants (non-experts), ranging between the ages of 18 and 40. In total, five different experiments were performed, the results of which are summarized in Table 1. We initially tested the capability of the presented platform through a game consisting of 5055 images, of which 471 were of infected RBCs and 4584 were of healthy RBCs (see Table 1). Additionally, 1266 (103 positives and 1163 negatives) RBC images were embedded as control images within the same game such that each gamer had to go through 6321 RBC images. The combined accuracy of the gamer diagnoses was 99%, with sensitivity (SE) of 95.1% and specificity (SP) of 99.4%. The positive predictive value (PPV) and negative predictive value (NPV) were also quite high at designer.
94.3% and 99.5% respectively (for definitions of SE, SP, PPV, and NPV refer to Table S1).

In addition to the gaming and the crowd-sourcing platform described earlier, we also developed an automated computer vision-based algorithm to detect the presence of malaria parasites (refer to Text S1–Section III and Figure S1 for details of implementation). In doing so our aim was to ultimately create a hybrid system such that machine vision and human vision can be coupled to each other, creating a more efficient and accurate biomedical diagnostics platform. For this purpose, independent of the human crowd, we next tested the automated diagnosis performance of our machine-vision algorithm, which was trained on 1266 RBC images (same as the control images used in experiment #1) and was tested on a total of 5035 unique RBC images (471 positives and 4584 negatives – see Table 1). This algorithm was able to achieve an overall accuracy of 96.3%, with SE-SP of 69.6%–99.0%, and PPV-NPV of 87.7%–96.9%. In terms of performance, our gamer crowd did better than this machine algorithm as summarized in Table 1. However, we should note that with an even larger training dataset (containing e.g., >10,000 RBC images) and more advanced classifiers, it may be possible to significantly improve the performance of our automated algorithm. This feat may be achieved through the coupling of statistical learning and crowd-sourcing into a hybrid model as illustrated in Figure 4, where a feedback exists between the gamers and the automated algorithm, resulting in a hybrid system as illustrated in Figure 4, where a feedback exists between the gamers and the automated algorithm, yielding an ever-enlarging training dataset as more games are played. This uni-directional feedback loop has the effect of labelling more and more images as training data for the automated algorithm, potentially leaving only the most difficult ones to be labelled by human gamers.

Following this initial comparison between human vision and machine vision for identification of malaria infected RBCs, to assess the viability of the above discussed hybrid diagnosis methodology, we conducted another test (experiments #3 & #4 in Table 1), where among all the RBC images characterized using our machine-vision algorithm, we extracted the ones with a diagnosis confidence level that is less than 30% of the maximum achieved confidence level, i.e. a total of 459 RBC images that were relatively difficult to diagnose were extracted. The training dataset (1266 RBC images that were used to train our machine algorithm, which also served as the control images of experiment #1) were then mixed with these “difficult-to-diagnose” 459 RBC images and were used to form a new game that is crowd-sourced to 27 human gamers. This new game (experiment #3) yielded an accuracy of 95.4%, with SE-SP at 97.8%–91.9%, and PPV-NPV at 94.7%–96.6% on these 459 difficult-to-diagnose RBC images. Next, we merged the results from the crowd-sourced game (experiment #3) and our machine algorithm (experiment #2) to arrive at an overall accuracy of 98.5%, with SE-SP of 89.4%–99.4% and PPV-NPV of 94.2%–98.9% (see experiment #4, Table 1). Thus, in this hybrid case we were able to increase the specificity and positive predictive value by 20% and 7%, respectively, and achieved a performance comparable to that of a completely human-labelled system (experiment #1), but with only 10% of the number of cells actually being labelled by humans. This significantly increases the efficiency of the presented gaming platform such that the innate visual and pattern-recognition abilities of the human crowd/gamers is put to much better use by only focusing on the “difficult-to-diagnose” images through the hybrid system (Figure 4).

In our next experiment (# 5) we increased the number of infected RBC images in the game by three-fold to simulate a scaled up version of the gaming platform. A total of 7829 unique RBC images were incorporated into the game, of which 784 were taken as control images that were repeatedly inserted into the game for a total of 2349 times. As a result, each gamer would go through 9394 RBC images, a quarter of which (2349) are known control images. Within the remaining 7043 test RBC images, there were 1549 (22%) positive images and 5496 negative images, which were all treated as unknown images to be diagnosed by the human crowd at the single cell level. The same ratio of positive to negative images was also chosen for the control RBC images in the game to eliminate any unfair estimation biases that may result from differing distributions. Completing this game (i.e. experiment # 5) took on average less than one hour for each gamer, and we can see in Table 1 that the accuracy of the overall human crowd (non-professionals) is within 1.25% of the diagnostic decisions made by the infectious disease expert. This experiment yielded an SE of 97.8% and an SP of 99.1%. The PPV was 96.7% and the NPV was 99.4%. The performance results of the individual players and their combined performances are shown in Figures S2 and S3.

Based on experiment #3, Figure 5 summarizes “the effect of the crowd” on diagnosis accuracy and sensitivity, i.e., how the overall performance of the crowd’s diagnosis is improved as more gamers are added to the system. We can see significant boosts in the sensitivity (i.e., the true positive rate) as diagnosis results from more gamers are added into the system. This is quite important as one of the major challenges in malaria diagnosis in sub-Saharan Africa is the unacceptably high false-positive rate, reaching ~60% of the reported cases [22]. Our overall diagnosis accuracy also steadily improves as more gamers are added as shown in Figure 5. This crowd effect may seem like a deviation from the traditional benefits of crowd-sourcing, in that multiple players are inaccurately solving the whole puzzle and then their results are combined to yield a more accurate solution. However, we should also note that cell images from a single blood smear slide can be broken up into multiple batches, where each batch is crowd-sourced to a group of players. In other words, each unique group of players will focus on one common batch of cell images, and in the end the diagnosis results will be combined once at the group level to boost the accuracies for each cell, and again at the slide level to make a correct overall diagnosis per patient. Therefore, the contribution of the crowd is twofold. First, it allows for the analysis problem to be broken up into smaller batches, and second, the analysis of the same batch by multiple individuals from the crowd allows for significantly higher overall diagnosis accuracies.

We should emphasise that throughout the manuscript we discuss diagnosis results for “individual” RBCs, not for patients. In reality, malaria diagnosis using a blood smear sample corresponding to a patient is a relatively easier task compared to single cell diagnosis since a thin blood smear for each patient sample already contains thousands of RBCs on it. Therefore statistical errors in the parasite recognition task could be partially hidden if the diagnostics decisions are made on a per blood-smear slide basis. To better demonstrate the proof of concept of our gaming based crowd-sourcing approach we aimed for the diagnosis of individual RBCs, rather than patients. Since any given patient’s blood smear slide will be digitally divided into smaller images (containing e.g., a handful of RBCs per image), and >1,000 RBC images per patient will be distributed to the crowd, we expect much higher levels of accuracy and sensitivity for diagnosis of individual patients. Furthermore, our single-cell-diagnosis-based gaming approach could also be very useful to estimate the parasitemia rate of patients which can be quite important and valuable for monitoring the treatment of malaria patients.

We should also emphasise that the work presented in this paper is a proof of concept and not the complete envisioned system, with potentially thousands of gamers and many patient slides to be diagnosed, which is left as future work. In addition to generating remote biomedical diagnosis through engaging games, the
presented platform can serve as an information hub for the global healthcare community as summarized in Figure 1. This digital hub will allow for the creation of very large databases of microscopic images that can be used for e.g., the purposes of training and fine tuning automated computer vision algorithms. It can also serve as an analysis tool for health-care policy makers toward e.g., better management and/or prevention of pandemics.

Next, we would like to brieﬂy discuss regulatory and practical issues that need to be addressed for deployment of the presented gaming and crowd-sourcing-based diagnosis and teledmedicine platform. As a potential future expansion of the platform, incentives (e.g., monetary ones) can be used to recruit health-care professionals who are trained and educated to diagnose such biomedical conditions, making them part of our gamer crowd. In such a scenario, one can envision the gaming platform to serve as an intelligent teledmedicine backbone that helps the sharing of medical resources through e.g., remote diagnosis and centralised data collection/processing. In other words, it would be a platform whereby the diagnosis can take place by professionals far away from the point-of-care. At the same time, it also enables the resolution of possible conﬂicting diagnostics decisions among medical experts, potentially improving the diagnostics outcome.

For this potentially highly trained crowd of “professional” gamers, the ﬁnal decisions made through the crowd can be used for direct treatment of the patient (without the need for regulatory approval). Furthermore, since these are trained medical professionals, the number of gamblers assigned to an image that is waiting to be diagnosed can be signiﬁcantly lower as compared to the case where “non-professional” gamers are assigned to the same image. On the other hand, if an image is diagnosed by entirely non-professional gamers, the result of the diagnosis can still be very useful to reduce the workload of health-care professionals located at point-of-care ofﬁces or clinics where the raw images were acquired. In the case of malaria diagnosis, this is especially relevant since the health-care professional is required to look at >1,000 RBC images for accurate diagnosis. Hence even a non-professional crowd’s diagnostics decisions could be highly valuable in guiding the local medical expert through the examination of a malaria slide, such that the most relevant RBC images are quickly screened ﬁrst, eliminating the need for conducting a manual random scan for rare parasite signatures.

Finally, the proposed methodology can be expanded to include a ‘training platform’. Assuming the expansion of this crowd-sourced diagnosticians platform and the generation of large image databases with correct diagnostics labels, software can be created to make use of such databases to assist in the training of medical professionals. Through such software, medical students and/or trainees can spend time looking at thousands of images, attempting diagnosis, and getting real-time feedback on their performances. Based on the concepts described in this paper, we also envision this platform to expand to other micro-analysis and diagnostics needs where biomedical images need to be examined by experts.

Supporting Information

Figure S1 Local Colour Peak Histograms (LCPH). For every window block, a colour histogram is calculated. The dominant pair of colours is used to compute an index (e.g., with 5 bins, there are a total of 10 different index values). A histogram of all index values is computed and used as part of the feature vector. In addition to colour-based features, we also used a number of more basic image features such as mean, variance, and gradient magnitude histograms to form our full feature vectors.

Figure S2 Individual performance results for experiment 5 of the Main Text. A total of 20 individual gamers played this game that consisted of 7045 test cell images with 1549 cells infected and 5496 cells healthy. We use a maximum a posterior probability (MAP) based approach to combine the results of multiple gamers and obtain higher performance levels. In order to show the worst case scenario, the worst N performances are combined in each MAP estimation.

Figure S3 Combined performance of gamers using the described MAP approach. We observe that if we combine all the available gamer data, we can determine the best gamer relative to others regardless of the ground truth data, lending itself to a scoring and ranking mechanism for gamers.

Table S1 Definition of acronyms used in the manuscript.

Text S1 This supporting text describes the mathematical and algorithmic details of the proposed framework.

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Author Contributions

Wrote the paper: SM AO. Designed and developed the theoretical framework and algorithms; contributed to the overall design of the games and related infrastructure; conducted the analysis and interpretation of the data and results: SM. Contributed to the development and implementation of the games and related infrastructure: SD SF FY. Contributed to the acquisition and compilation of microscopic images: US OY SP. Contributed to the medical implementation of the presented platform: KN. Conceived the idea, supervised the project, and contributed to the overall design of the games, related infrastructure and theory as well as to the analysis and interpretation of the data and the results: AO.
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SUPPLEMENTARY TEXT

I- Definitions of acronyms used in the manuscript

The acronyms used in this manuscript are defined in Table S1.

II- Theoretical Framework for Crowd-sourced Malaria Diagnosis Game

In this section we will describe the theoretical framework with which we have approached the problem of fusing the data from multiple individuals playing the game. In doing so, we used communications and probability theory based methods.

*Binary Channel Model for Gamers Diagnosing Red Blood Cells:* Since we are attempting to combine decisions that are received from many gamers, we will be feeding them the same set of images to label. Therefore, there is a single sequence of images to be labelled, and each gamer will output a decision sequence. Ideally, the output of each gamer yields the correct diagnostic labels for the blood cell images. Given that each image either corresponds to a healthy cell or an infected cell, we can use binary labels to identify them: 0 for healthy and 1 for infected. Recasting our setup as a communications system, our server will act as a broadcaster of a binary sequence and each gamer will act as a noisy Binary Channel\(^1\), retransmitting the symbols back along with some errors (see Figure 3 of Main Text for further details). Note that since the gamers may not necessarily make mistakes symmetrically, the probability of a gamer mislabelling a healthy cell may be different from that of mislabelling an infected cell.

*System Overview:* We model the framework of our games as a communications system consisting of a broadcast unit, multiple repeaters, and a receiver/decoder as shown in Figure 3 of the Main Text. In the ideal scenario, the repeaters (i.e., the gamers) would simply receive a set of incoming symbols (images to be diagnosed) from the broadcast unit, and transmit them to the receiver/decoder block, which in turn combines and stores the decisions (image labels).

*Broadcast Unit - The Source of the Microscopic Images:* We can model the source of our microscopic images as a broadcast unit. In this analogy, the cell images are essentially treated as binary symbols. We assume the equivalency of the two image classes infected and healthy with binary symbols 1 and 0 respectively.

The sequence of symbols \(x_1, \ldots, x_N\) is broadcast to \(M\) repeater units (i.e., the gamers within the crowd) that can be viewed as \(M\) parallel noisy channels (see Figure 3 of Main Text). To be able to decode the outputs of these channels reliably, it is necessary to *learn* the channels adaptively. As such, it is crucial to embed some known symbols (i.e., *control images*) in the
output of the broadcast unit. Knowing the binary value of certain symbols/images at specific times, we can learn the conditional probabilities \( p_j(y_i|x_i) \) as more symbols are transmitted by the broadcast unit and passed through the repeaters/gamers.

Additionally, an encoder unit can also be placed after the broadcast unit to increase the redundancy of the transmission, and allow for error correction at the decoder.

**Gamers as Repeaters in the Communication System:** Each gamer is modelled as an independent repeater that behaves as a binary channel (see Figure 3 of the Main Text). We define the error probabilities using the notation \( p_j(y_i|x_i) \), corresponding to the probability that the \( j^{th} \) user will output the symbol \( y_i \) when observing the symbol \( x_i \) (i.e. \( i^{th} \) image). In general, the error probabilities are asymmetric, i.e., \( p(y = x|x) \) is not the same for different values of \( x \). However, it is difficult to accurately estimate this asymmetric probability in our games due to the imbalance in the existence of positive and negative training data (*which is true not only for malaria diagnosis but also for various biomedical image analysis/diagnosis problems such that disease signatures are relatively rare compared to healthy data*) which causes a bias towards better estimating the error probabilities when \( x = 0 \) (healthy case). In addition to this, another practical limitation is due to the general infeasibility of embedding large numbers of training images within the game. It is therefore more straightforward to estimate a simpler bit-flip probability, assuming a Binary Symmetric Channel (BSC)\(^1\). It was observed in our experiments that a BSC model performs better than an asymmetric model. This observation stems from the fact that there is an inherent imbalance in the number of positive and negative image samples. Since we are using a limited number of control images to estimate the probability densities of the gamers’ performances, this imbalance in the control data translates to having a small number positive samples for accurate estimation of \( p_{11} \) and \( p_{10} \).

**Error Control Coding (ECC) in Malaria Diagnosis Games:** Similar to traditional communications systems, our broadcast unit can also include an encoder to increase the information redundancy prior to transmission to the repeaters/gamers. Given that the symbols being transmitted by the broadcast unit are not known *a priori*, the only appropriate coding scheme is the repetition code, where each symbol is repeated for an odd number of times prior to transmission. At the decoder, a majority vote is taken on the channel outputs.

Leaving the communications system analogy, an ECC amounts to asking the gamer to play each image an odd number of times, and the most frequently assigned label is taken to be his or her answer for that particular image. This can also be interpreted as the gamer’s confidence in the given diagnosis response. In other words, if the gamer is absolutely sure of a particular diagnosis for an image, then he or she will choose the same label on every observation. However, if the image is difficult to diagnose, then there is the chance that the gamer would not be consistent in making a decision, thus producing a lower confidence level.
**Decoder for fusing Gamers’ Decisions toward Malaria Diagnosis:** In designing the decoder for our gaming platform, we take a Maximum a Posteriori Probability (MAP) approach. Suppose that we have \(N\) symbols/images \(x_1, \ldots, x_N\) being broadcast and relayed through \(M + 1\) repeaters/gamers and are received by the decoder. Also assume that we have estimates of the repeater/gamer error probabilities \(p_j(y_i|x_i)\) for the \(j^{th}\) repeater and the \(i^{th}\) symbol. We then would like to estimate \(x_i\) given all the repeater outputs \(y_i^0, \ldots, y_i^M\) (see Figure 3 of the Main Text). For a particular transmitted symbol \(x^*\) we have:

\[
p(x_i = x^*|y_i) = \frac{p(y_i|x_i = x^*) \cdot p(x_i = x^*)}{p(y_i)} = \prod_{j=0}^{M} p(y_j^i|x_i = x^*) \cdot \frac{p(x_i = x^*)}{\sum_{l=0}^{1} [p(x_i = l) \prod_{k=0}^{M} p(y_k^i|x_i = l)]}
\]

We can then have:

\[
\log p(x_i = x^*|y_i) = \sum_{j=0}^{M} \log p(y_j^i|x_i = x^*) + \log p(x_i = x^*) - \log \left[ \sum_{l=0}^{1} p(x_i = l) \prod_{k=0}^{M} p(y_k^i|x_i = l) \right]
\]

The value of \(z_i\) is taken to be \(x^*\) that maximizes the above posterior. Therefore, we have:

\[
z_i = \arg \max_{l \in \{0, 1\}} \left[ \varphi_l + \sum_{j=0}^{M} \log p(y_j^i|x_i = l) \right], \quad \varphi \equiv \log p(x_i = l)
\]

**Automated Malaria Diagnosis using Machine Vision:** In addition to the gaming and the crowdsourcing platform described earlier, we also developed an automated machine-learning-based algorithm to detect the presence of malaria parasites (i.e., \(P. falciparum\)). In doing so our aim was to ultimately create a hybrid system such that machine vision and human vision can be coupled to each other to create a more efficient and accurate diagnostics platform as illustrated in Figure 4 of the Main Text.

Toward this end, instead of developing a very specific machine vision algorithm that is only applicable to malaria diagnosis we opted to create one that could potentially be also applied to...
other infections that may have different visual properties. As such we developed a machine learning algorithm that extracts local colour features from cell images and feeds them into a classifier. In training the classifier, we used a small subset of the cell images in our dataset (same images that are embedded as ‘control images’ in the game) as the training set. We then used a variation of Adaptive Boosting $^{2,3}$ to create an ensemble of relatively weak classifiers that together produce accurate classification results as detailed in our Results section of the Main Text. For further details of our machine-learning based malaria diagnosis algorithm, please refer to the Section III of this Supplementary Text.

**Hybrid (Machine + Human) Diagnosis Framework:** As briefly pointed out earlier, the two discussed diagnosis methodologies (crowd-sourced gaming and machine vision) naturally lend themselves to a third hybrid diagnostics platform (see Figure 4 of the Main Text) where the images are initially processed by automated machine vision algorithms with a measure of confidence for each diagnostics decision. Images for which the machine decisions are not above a certain threshold (e.g., T) are then passed on to the human gamers to be re-diagnosed by the crowd. As demonstrated in our Experiment #4 of the Main Text, this can significantly increase the efficiency of the human gamers, since only those images that are deemed as “difficult-to-diagnose” by the machine vision algorithm are passed to the gamers, allowing the same number of gamers to assist in e.g., the diagnosis of more blood smears and patients per unit time.

**III- Details of our Automated Recognition Algorithm for Malaria Parasites**

In this manuscript our focus with regards to machine learning algorithms was to develop a general methodology such that it can also be applied to other parasites or disease signatures beyond malaria with small modifications in the visual features used for analysis. Therefore, in addition to the gaming based crowd-sourcing platform, we also developed an automated machine learning algorithm to detect the presence of malaria parasites, which helps us to compare our gaming platform’s diagnosis accuracy to state-of-the-art machine vision algorithms as well as to ultimately create a hybrid system such that machine vision and human vision can be coupled to each other to create a more efficient and accurate diagnostic platform as illustrated in experiment #4 of the Main Text. In the next sub-sections we will go over some details of our algorithm.

**Selection of Image Features for Recognition of Malaria Parasites:** On a thin blood smear, malaria parasites are usually stained with Giemsa stain and are digitally imaged under a
brightfield microscope using a 100X oil-immersion objective lens. Giemsa stains the nuclear material in the parasite with a blue tint, and does not affect the RBC morphology.

In general, there are two possible approaches in extracting features from the acquired microscope images for the purpose of building a digital classifier. One can either attempt to extract very specific, hand-coded features or try to learn discriminative features from a large set of training examples. The advantage of using hard-coded features is that we can use prior knowledge of the physical/structural properties of the parasites. For example, we can look for “ring-shaped objects” within the RBC image as an indicator for the existence of the parasite. The advantage of using such features is that they lend themselves to very fast implementations and make the job of the classifier much easier. On the other hand, they are difficult to design, and are in general very inflexible to variations in sample preparation or illumination/recording conditions. For example, custom-designed malaria feature-sets that use shape and colour may not be easily modified to detect a different parasite type.

In contrast to hand-coded features, ‘learned’ features can be very generic and easily modified and applied to similar detection problems. They also take less time and effort to design for a particular problem, and put most of the burden of classification on the classifier itself. In this work, we preferred the second approach (i.e., learned features) and used a set of generic colour-based features for training a classifier to discriminate between RBC images that contain *P. falciparum* and those which do not.

We used two types of histogram-based features as input to our classifier. The first is a simple colour histogram of the image in grayscale. This is a feature that carries information about the general distribution of image values. We used a second, more complicated colour feature which we dubbed *Local Colour Peak Histograms* (LCPH). The LCPH for an image is formed by first generating highly quantized colour histograms in the Hue-Saturation space over local windows. For each window the two most occurring histogram bins are found. Any given pair of bins corresponds to a particular index value. In other words, an index can be assigned to the occurrence of each pair of values in the histogram, and this is the value that is recorded for each local window in the image. A second histogram is generated over the recorded indexes of all the local windows and used as the LCPH features. This feature essentially measures the relative occurrences of various colour pair co-occurrences throughout the image (see Error! Reference source not found.).

In addition to the above mentioned colour-based features, we used a number of more basic image features such as mean, variance, and gradient magnitude histograms to form our final feature vectors.
**Classifier Training:** In training our malaria classifier, we used a variation of Adaptive Boosting and trained many weak decision-tree classifiers that together can produce a very strong classifier. In classical Adaptive Boosting, the overall classifier is tested on the complete training dataset at each iteration. Data points that are not correctly classified are then assigned larger weights for the next classifier to be trained. This is where we deviate from the classical algorithm in that instead of re-weighting the full training set and training a new classifier, we use the weights to probabilistically select a small subset of the training data for the next weak classifier to be trained. This allows for completely disjoint training data for the weak classifiers and results in very fast convergence of the boosted classifier. Refer to Algorithm S1 shown below for further details of our implementation.

**Algorithm S1** Summary of our Adaptive Boosting Algorithm. The total number of training points is fixed for each weak classifier, i.e., for each weak classifier a total of $n_S$ training vectors are chosen based on the weights $W_k(i)$.

**IV- Detailed analysis of gamers’ performance**

Detailed analysis of our largest gaming experiment (#5 - see Table 1 of the Main Text) is shown in Figures S2 and S3.

**V- Sources of our Microscopic Data**

The RBC images used in this work were obtained from multiple sources, which is quite important as it mimics a real-life scenario where various different optical microscopes located
at e.g., point-of-care offices and malaria clinics would be used in our gaming based diagnosis platform. The bulk of the images were captured using an in-house optical microscope with a 100X oil-immersion objective lens (numerical aperture: 1.25). Roughly 5% of the positive images and 10% of the negative images were also obtained from Center for Disease Control (CDC) database. Figure S4 shows representative images demonstrating the colour, illumination and background variability observed among these RBC images that were used in our crowd-sourced games.

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| Term                      | Acronym | Definition                                                   |
|---------------------------|---------|--------------------------------------------------------------|
| True Positive             | TP      | Number of correctly labelled positive samples                |
| False Positive            | FP      | Number of negative samples incorrectly labelled as positive |
| True Negative             | TN      | Number of correctly labelled negative samples                |
| False Negative            | FN      | Number of positive samples incorrectly labelled as negative  |
| Accuracy                  | ACC     | $\frac{TP + TN}{TP + TN + FP + FN}$                        |
| Sensitivity or True Positive Rate | SE or TPR | $\frac{TP}{TP + FN}$                                      |
| False Positive Rate       | FPR     | $\frac{FP}{TN + FP}$                                       |
| Specificity or True Negative Rate | SP or TNR | $\frac{TN}{TN + FP}$                                      |
| Positive Predictive Value | PPV     | $\frac{TP}{TP + FP}$                                       |
| Negative Predictive Value | NPV     | $\frac{TN}{TN + FN}$                                       |
| Image Source                                      | Positive Sample | Negative Sample |
|--------------------------------------------------|-----------------|-----------------|
| Center for Disease Control (CDC)                 | ![Images]       | ![Images]       |
| CDC                                              | ![Images]       | ![Images]       |
| CDC                                              | ![Images]       | ![Images]       |
| Cultured Smears captured using Meiji MX w/ 100x/1.25 NA oil immersion objective | ![Images]       | ![Images]       |