

# Learning from Crowds with Variational Gaussian Processes

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## Abstract

Solving a supervised learning problem requires to label a training set. This task is traditionally performed by an expert, who provides a label for each sample. The proliferation of social web services (e.g., Amazon Mechanical Turk) has introduced an alternative crowdsourcing approach. Anybody with a computer can register in one of these services and label, either partially or completely, a dataset. The effort of labeling is then shared between a great number of annotators. However, this approach introduces scientifically challenging problems such as combining the unknown expertise of the annotators, handling disagreements on the annotated samples, or detecting the existence of spammer and adversarial annotators. All these problems require probabilistic sound solutions which go beyond the naive use of majority voting plus classical classification methods. In this work we introduce a new crowdsourcing model and inference procedure which trains a Gaussian Process classifier using the noisy labels provided by the annotators. Variational Bayes inference is used to estimate all unknowns. The

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proposed model can predict the class of new samples and assess the expertise of the involved annotators. Moreover, the Bayesian treatment allows for a solid uncertainty quantification. Since when predicting the class of a new sample we might have access to some annotations for it, we also show how our method can naturally incorporate this additional information. A comprehensive experimental section evaluates the proposed method with synthetic and real experiments, showing that it consistently outperforms other state-of-the-art crowdsourcing approaches.

*Keywords:* Crowdsourcing, Classification, Gaussian Processes, Bayesian Modeling, Variational Inference.

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## 1. Introduction

The main goal in supervised learning is to find a mapping that predicts labels from features [1, 2, 3]. Most of the works in supervised learning assume that the training samples have been labeled with no errors by an expert [4, 5].  
5 However, the recent advent of social web services has introduced a new approach to address the labeling problem. The term *crowdsourcing* was coined in 2006 by J. Howe [6] to describe “*the act of taking a job traditionally performed by a designated agent (usually an employee) and outsourcing it to an undefined generally large group of people in the form of an open call*”.

10 In the last decade, many crowdsourcing services have proliferated in the Internet, where a dataset can be published and millions of people around the world can provide labels in exchange for a reward [7]. Amazon Mechanical Turk ([www.amt.com](http://www.amt.com)), Galaxy Zoo ([www.galaxyzoo.org](http://www.galaxyzoo.org)), Zooniverse ([www.zooniverse.org](http://www.zooniverse.org)), Crowdflower ([www.crowdflower.com](http://www.crowdflower.com)) or Clickworker ([www.clickworker.com](http://www.clickworker.com))  
15 are among the most popular ones. Due to the great number of potential annotators, large data sets can be labeled in a very short time. However, this approach introduces new challenging problems: combining the unknown expertise of annotators, dealing with disagreements on the annotated samples, or detecting the existence of spammer and adversarial annotators [7].

20 The first paper on crowdsourcing dates back to 1979 [8]. Early contributions attempted to estimate the underlying true labels and the reliability of the annotators, but were not conceived to learn a classifier. This idea was explored by Raykar *et al.* [9], who proposed to jointly estimate the coefficients of a logistic regression (LR) classifier and the annotators’ expertise. The latter is modeled  
25 through the *sensitivity* and *specificity* concepts, which refer to the accuracy of the annotator when labelling instances from each class. Yan *et al.* [10] (see also the subsequent journal version [11]), introduced a crowdsourcing classifier (also based on LR) which considers a feature-dependent model for the annotators’ expertise. The main limitation of these two approaches is the simple LR classification model, which can only deal with linearly separable data. Rodrigues *et al.* [12] overcame this problem by introducing a crowdsourcing classifier based on Gaussian Processes (GP) [13, 14, 15]. GP theory makes use of the so-called “kernel trick” [1, Chapter 6] to model complex classification problems where the decision boundary may be non-linear. Expectation Propagation (EP) [13, Section 3.6] is used as inference procedure in [12]. To the best of our knowledge,  
30 this is the most recent general-purpose probabilistic crowdsourcing approach (see also [16, Section 2.2].)

Nowadays, crowdsourcing is a really active and promising research field, in which these general-purpose crowdsourcing methods are being tailored to a  
40 wide range of relevant problems (see the recent survey [7] and related works [17, 18]). Crowdsourcing is being applied to modern areas such as ecological monitoring and conservation [19], plant phenotyping [20, 21], remote sensing [22], mitosis detection in breast cancer histology images [23], topic modeling from crowds [16], and detection of glitches in signals acquired by the laureate  
45 Laser Interferometer Gravitational-Wave Observatory (LIGO) [24]. Moreover, there exist some recent attempts to combine crowdsourcing with Deep Learning approaches [23, 25], and new challenges, such as the optimal expert validation of the crowdsourced labels [26], are emerging.

In this work we address the crowdsourcing classification problem. As in [12],  
50 the true underlying training labels are modeled as latent variables by means of

a GP. A sensitivity-specificity model is used for the annotators (as in [9] and [12]). However, there exist two main differences with [12]: 1) we use Variational Bayes inference (VB) to estimate all unknowns (instead of EP), and 2) we model sensitivity and specificity as stochastic variables (instead of point parameters).  
55 Several reasons motivated our choice of Variational inference. First, it is well-known that the EP iterative procedure does not guarantee convergence, and it may not be able to capture complex posterior distributions (e.g., multi-modal) [1, Section 10.7]. Second, as it will become clear in the experiments, the EP inference is usually slower in practice (which, in fact, has led to the introduction  
60 of some strategies to optimize it [27]). A thorough experimentation (including comparisons with the aforementioned approaches in [9], [10] and [12] among others) will show that the proposed ideas can contribute to advance the current state-of-the-art in crowdsourcing classification. Moreover, the proposed model naturally lends itself to the integration of annotations that may have been pro-  
65 vided for test instances in the prediction of their true class. The experiments will show that, if test annotations are available, this hybrid human-machine prediction is significantly more accurate than the one produced by either the machine or the annotators alone. To the best of our knowledge, this extension had not been addressed in any previous work.

70 This paper gathers together, clarifies, and significantly extends the ideas in our two conference contributions [28, 29]. The main novelties are: first, sensitivity and specificity are treated as stochastic variables (they are estimated through non-degenerate posterior distributions instead of point estimates). This allows for a better uncertainty quantification and, thus, an enhancement in the  
75 experimental results. Second, we show how our model can naturally integrate in the prediction annotations that may have been provided for test instances. If there are no such annotations, the new predictive distribution recovers the old one. Third, the experiments are exhaustively extended in several ways: the new methodology to integrate test set annotations is evaluated, a new type of data  
80 popular in crowdsourcing is introduced (*semi-synthetic* data), the computational cost is assessed, and the annotators' expertise estimations are reported in all

experiments. Fourth, the experimental section does not restrict itself to the performance of the proposed method, but also examines the behavior of other state-of-the-art approaches that it is compared against. Thus, it can be useful  
85 as a brief experimental review of the main current crowdsourcing methods.

The rest of the paper is organized as follows. To facilitate the reading of the paper, an exhaustive glossary of all the symbols used in this work is included in Table 1. Section 2 presents the proposed probabilistic crowdsourcing model based on GP. The VB inference procedure is described in Section 3. The process  
90 to classify new samples (including the case when there are test annotations available) is described in Section 4. A comprehensive experimental validation is presented in Section 5. Section 6 concludes the paper and provides some future outlook.

## 2. Bayesian Modeling

Let  $\mathbf{X} = [\mathbf{x}_1, \dots, \mathbf{x}_N]^T \in \mathbb{R}^{N \times D}$  be a training set of  $N$   $D$ -dimensional sam-  
95 ples, with unknown labels  $\mathbf{z} = (z_1, \dots, z_N)^T \in \{0, 1\}^N$ . Let us assume there are  $R$  different annotators. Let  $R_n \subseteq \{1, \dots, R\}$  denote the subset of annotators who labeled the  $n$ -th sample, and  $N_r \subseteq \{1, \dots, N\}$  the subset of samples labeled by the  $r$ -th annotator. Finally,  $\mathbf{Y} = \{y_n^r \in \{0, 1\} \mid n = 1, \dots, N; r \in R_n\}$  is the  
100 set of labels provided by the  $R$  annotators.

Gaussian Processes (GP) model the relationship between samples  $\mathbf{X}$  and the corresponding unknown true labels  $\mathbf{z}$  in two steps. First, a set of latent variables  $\mathbf{f} = [f_1, \dots, f_N]^T$  following a joint Gaussian distribution  $p(\mathbf{f}|\mathbf{\Omega}) = \mathcal{N}(\mathbf{f}|\mathbf{0}, \mathbf{K})$  is introduced. The kernel matrix  $\mathbf{K} = [k(\mathbf{x}_n, \mathbf{x}_m|\mathbf{\Omega})]_{nm}$  is computed with  
105 the kernel function  $k$ , which defines an inner product in a (possibly infinite-dimensional) transformed space [1, Chapter 6]. Intuitively, the correlation between each pair of entries of  $\mathbf{f}$  is calculated in a transformed space of the original feature space, which allows GP to estimate non-linear decision boundaries. In this work we use the well-known squared exponential (SE) kernel  
110  $k(\mathbf{x}_n, \mathbf{x}_m) = \gamma \cdot \exp(-\|\mathbf{x}_n - \mathbf{x}_m\|^2 / (2l^2))$ , although other kernels could be used.

Symbol	Description
$\mathcal{C}_0$ and $\mathcal{C}_1$	Classes 0 and 1, respectively.
$N$	Number of samples in the training set.
$D$	Dimension of the feature space.
$R$	Number of annotators who provided crowdsourcing labels.
$R_n \subseteq \{1, \dots, R\}$	Subset of annotators who labeled the $n$ -th sample.
$N_r \subseteq \{1, \dots, N\}$	Subset of samples labeled by the $r$ -th annotator.
$\mathbf{X} \in \mathbb{R}^{N \times D}$	Matrix containing all the samples in the training set.
$\mathbf{x}_n \in \mathbb{R}^D$	$n$ -th sample of the training set.
$\mathbf{x}_* \in \mathbb{R}^D$	New sample whose class is predicted by the proposed method.
$\mathbf{Y}$	Set of annotations provided by the $R$ annotators.
$y_n^r \in \{0, 1\}$	Label provided by the $r$ -th annotator for the $n$ -th sample.
$\mathbf{y}_* = \{y_*^r : r \in R_*\}$	Annotations provided for the new sample $\mathbf{x}_*$ .
$\mathbf{z} \in \{0, 1\}^N$	Underlying real labels for the training set instances.
$z_n \in \{0, 1\}$	Underlying real label for the $n$ -th sample.
$z_* \in \{0, 1\}$	Underlying real label for $\mathbf{x}_*$ .
$\boldsymbol{\alpha} = [\alpha_1, \dots, \alpha_R]$	Sensitivity of each annotator.
$\boldsymbol{\beta} = [\beta_1, \dots, \beta_R]$	Specificity of each annotator.
$a_0^\alpha, b_0^\alpha, a_0^\beta$ and $b_0^\beta$	Hyperparameters for $\alpha$ and $\beta$ . (Default: All of them equal to 1.)
$\mathbf{f} = [f_1, \dots, f_N]^T$	GP modeling the relationship between $\mathbf{X}$ and $\mathbf{z}$ .
$f_* \in \mathbb{R}$	GP modeling the relationship between $\mathbf{x}_*$ and $z_*$ .
$k(\cdot, \cdot   \boldsymbol{\Omega})$	Kernel function depending on a set of parameters $\boldsymbol{\Omega}$ .
$\mathbf{K} \in \mathbb{R}^{N \times N}$	Covariance matrix of the prior distribution of $\mathbf{f}$ .
$\mathbf{h} \in \mathbb{R}^N$	Vector of prior covariances between $f_*$ and $f_1, \dots, f_n$ .
$c \in \mathbb{R}^+$	Prior variance of $f_*$ .
$\boldsymbol{\xi} = (\xi_1, \dots, \xi_N)^T$	Variational parameters to be estimated.
$\boldsymbol{\Lambda} \in \mathbb{R}^{N \times N}$	Diagonal matrix calculated from the variational parameters $\boldsymbol{\xi}$ .
$m_*$ and $s_*^2$	Mean and variance of the approximated posterior distribution of $f_*$ .
$\boldsymbol{\mu}_f$ and $\boldsymbol{\Sigma}_f$	Mean and covariance matrix of the posterior distribution of $\mathbf{f}$ .
$\delta \in [0, 1]$	Classification threshold.
$\Theta = \{\mathbf{f}, \boldsymbol{\alpha}, \boldsymbol{\beta}, \boldsymbol{\Omega}\}$	Subset of unknown variables of the model.
$\bar{\Theta} = \{\mathbf{z}, \mathbf{f}, \boldsymbol{\alpha}, \boldsymbol{\beta}, \boldsymbol{\Omega}\}$	Set of the unknown variables of the model.
$\bar{\Theta}_\theta = \bar{\Theta} \setminus \theta$	Set $\bar{\Theta}$ minus the element $\theta \in \bar{\Theta}$ .
$\sigma(\cdot)$ and $\psi(\cdot)$	Sigmoid and Digamma functions, respectively.
$\text{KL}(\cdot    \cdot)$	Kullback-Leibler divergence.
$p(\cdot)$ and $q(\cdot)$	Probability distributions: Assumed known (p) and approximated (q).
$\mathbf{0}$ and $\mathbf{1}/2$	Vector with all the components equal to 0 and 1/2, respectively.

Table 1: A comprehensive glossary of all the symbols used in this work.

The kernel hyperparameters  $\boldsymbol{\Omega} = \{\gamma, l\}$  are called *variance* and *length-scale*, respectively.

The second step is to relate the latent variables  $\mathbf{f}$  to the unknown true labels  $\mathbf{z}$  using a product of Bernoulli distributions:

$$p(\mathbf{z}|\mathbf{f}) = \prod_{n=1}^N \sigma(f_n)^{z_n} (1 - \sigma(f_n))^{1-z_n} = \prod_{n=1}^N \left( \frac{1}{1 + e^{-f_n}} \right)^{z_n} \left( \frac{e^{-f_n}}{1 + e^{-f_n}} \right)^{1-z_n}, \quad (1)$$

115 where the sigmoid function  $\sigma$  maps  $\mathbb{R}$  into the interval  $(0, 1)$ . In other words, the sigmoid function assigns the probability of belonging to a class depending on the value of the real number  $f_n$ . When  $\mathbf{x}_n$  belongs to class 1 ( $z_n = 1$ ), only the first factor is considered and a large positive value is expected for  $f_n$ . When  $\mathbf{x}_n$  belongs to class 0 ( $z_n = 0$ ), only the second factor appears and a large  
 120 negative value is expected for  $f_n$ . Notice that, although a realization of a GP is a continuous real function on the feature space, the sigmoid function transforms it into a Bernoulli parameter. This is a natural generalization of logistic regression [30]. While logistic regression uses a linear combination of the components of  $\mathbf{x}_n$ , with linear weights to be estimated, GP uses a linear combination of features in a  
 125 transformed domain (this transformed domain depends on the kernel used) and denotes by  $f_n$  the corresponding linear combination. Moreover, the sigmoid function is an infinitely differentiable function, which allows VB to infer the posterior distribution of the latent variable  $\mathbf{f}$ .

The distributions  $p(\mathbf{f}|\boldsymbol{\Omega})$  and  $p(\mathbf{z}|\mathbf{f})$  define a standard GP classifier. Now we need to include the crowdsourcing labelling process in our model. Each annotator  $r$  is described by their *sensitivity*  $\alpha_r := p(y^r = 1|z = 1)$  and *specificity*  $\beta_r := p(y^r = 0|z = 0)$ . Intuitively,  $\alpha_r$  and  $\beta_r$  represent the reliability of the  $r$ -th annotator when labeling samples of class  $\mathcal{C}_1$  and  $\mathcal{C}_0$ , respectively. This model is the same as in [9, 12]. Assuming independence between annotators, we have the following product of Bernoulli distributions

$$p(\mathbf{Y}|\mathbf{z}, \boldsymbol{\alpha}, \boldsymbol{\beta}) = \prod_{r=1}^R \prod_{n \in N_r} \left[ \alpha_r^{y_n^r} (1 - \alpha_r)^{1-y_n^r} \right]^{z_n} \left[ (1 - \beta_r)^{y_n^r} \beta_r^{1-y_n^r} \right]^{1-z_n}, \quad (2)$$

where  $\boldsymbol{\alpha} = (\alpha_1, \dots, \alpha_R)$  and  $\boldsymbol{\beta} = (\beta_1, \dots, \beta_R)$ . Some observations are required  
 130 at this point. First, this sensitivity-specificity model allows for scenarios where

annotators might be non-experts. Second, *spammer* (resp. *adversarial*) annotators are those with  $\alpha_r$  and  $\beta_r$  values close to (resp. much lower than) 0.5. Third, notice that exchanging the role of  $z_n$  and  $1 - z_n$  in eq. (2) corresponds to exactly the same model but with sensitivities  $1 - \beta_r$  and specificities  $1 - \alpha_r$ .  
135 This means that, changing  $\boldsymbol{\alpha}$  and  $\boldsymbol{\beta}$  accordingly, a certain set of underlying true training labels and their opposite are equally plausible. In section 3 we provide an initialization of our algorithm that accounts for this ambiguity.

In this work, sensitivities and specificities are treated as stochastic variables, and Beta distributions are used as hyper-priors. This treatment allows us to introduce prior knowledge about these parameters and weigh more certain configurations of them, yielding more accurate results and a better uncertainty quantification of the model. The independence between annotators yields  $p(\boldsymbol{\alpha}) = \prod_{r=1}^R \text{Beta}(\alpha_r | a_0^\alpha, b_0^\alpha)$  and  $p(\boldsymbol{\beta}) = \prod_{r=1}^R \text{Beta}(\beta_r | a_0^\beta, b_0^\beta)$ , where we have removed the dependency on the parameters for simplicity. Recall that  $\text{Beta}(\omega | a, b) \propto \omega^{a-1}(1 - \omega)^{b-1}$  with mean  $\langle \omega \rangle = a/(a + b)$ . During inference, the following expectations will be required (see [1, Exercise 2.11])

$$\langle \log \omega \rangle = \psi(a) - \psi(a + b), \quad \langle \log(1 - \omega) \rangle = \psi(b) - \psi(a + b), \quad (3)$$

where  $\psi$  denotes the digamma function. The parameters  $a$  and  $b$  can be set to introduce prior knowledge about the expected values of  $\boldsymbol{\alpha}$  and  $\boldsymbol{\beta}$ , and our  
140 confidence on them. When no prior knowledge is available,  $a = b = 1$  produces uniform distributions. For instance, these hyper-priors on  $\boldsymbol{\alpha}$  and  $\boldsymbol{\beta}$  are useful to deal with annotators who only provide labels for samples in one of the classes (see [2] for more details about the so-called “*black swan paradox*”).

The proposed joint probabilistic model for the crowdsourcing problem is

$$p(\mathbf{Y}, \mathbf{z}, \mathbf{f}, \boldsymbol{\alpha}, \boldsymbol{\beta}, \boldsymbol{\Omega}) = p(\mathbf{Y}, \mathbf{z}, \boldsymbol{\Theta}) = p(\mathbf{Y} | \mathbf{z}, \boldsymbol{\alpha}, \boldsymbol{\beta}) p(\mathbf{z} | \mathbf{f}) p(\mathbf{f} | \boldsymbol{\Omega}) p(\boldsymbol{\alpha}) p(\boldsymbol{\beta}) p(\boldsymbol{\Omega}), \quad (4)$$

where  $\boldsymbol{\Theta} = \{\mathbf{f}, \boldsymbol{\alpha}, \boldsymbol{\beta}, \boldsymbol{\Omega}\}$ , and  $p(\boldsymbol{\Omega})$  is a flat prior on the kernel parameters  
145  $\boldsymbol{\Omega}$ . The probabilistic graphical model is depicted in Figure 1. Yellow nodes correspond to observed variables, namely, the set of features  $\mathbf{X}$  and the labels provided by the annotators  $\mathbf{Y}$  (discrete). The unknown variables, to be inferred

during training, are represented using blue nodes, namely, the real labels  $\mathbf{z}$  (discrete), the GP latent variable  $\mathbf{f}$  (continuous), the GP hyper-parameters  $\mathbf{\Omega}$  (continuous), and sensitivity and specificity  $\alpha, \beta$  (continuous).  
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Now that we have the full probabilistic modeling of our problem, let us briefly describe and explain in words its capabilities and limitations. First we utilize a GP (a prior on the set of functions defined over the feature space) which in combination with the sigmoid function is used to describe the real underlying classifier. Since we do not have access to the output of this classifier, the probability distributions of the labels provided by each annotator given the output of the true classifier is modeled using the sensitivity (when the true label is one) and specificity (when the true label is zero) parameters. These numbers quantify how close each annotator’s behavior is to the true classifier. Finally, any additional information on each annotator’s behavior can be included as prior information on the two aforementioned parameters. In summary, we are using a sound and robust to outliers probabilistic modeling of our crowdsourcing problem  
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One of the main limitations of the proposed framework is that the only supervised source of information consists in the labels provided by annotators, lacking some mechanism to introduce additional supervised knowledge which may prevail over the annotators. For instance, there may be some instances in the training set for which we know the real label instead of just (noisy) annotations, in which case we would rather rely on this more accurate knowledge. Second, a simple model has been considered for the annotators, based solely on their sensitivity and specificity. More complex (in particular, feature-dependent) behaviors could happen in real-world problems. For instance, there might be annotators who are much more skilled when labelling instances coming from a certain region of the feature space (because they have specialized in that type of instances), but are not that reliable in other regions. Third, there is the implicit assumption that all the annotators are not spammers. Notice that, for the scenario where all the annotators provide random labels for all the instances, there is no information to be able to infer the true decision boundaries, in which  
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case the proposed method cannot train an accurate classifier.

180 Having explained the model, let us now see how inference is carried out, what problems will be found when estimating the posterior distribution of all the unknowns given the labels provided by the annotators, and how variational inference can be used to solve all of them.

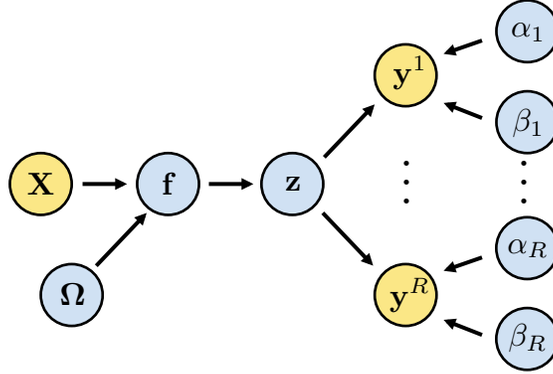


Figure 1: Probabilistic graphical representation for the proposed model. Yellow nodes denote observed variables, and blue nodes unknown variables (to be inferred during training).  $\mathbf{Y} = \{\mathbf{y}^1, \dots, \mathbf{y}^R\}$  and  $\mathbf{z}$  are discrete variables, whereas the rest are continuous.

### 3. Variational Bayes Inference

185 In Bayesian inference, the main goal is to find the posterior distribution  $p(\mathbf{z}, \Theta | \mathbf{Y}) = p(\mathbf{Y}, \mathbf{z}, \Theta) / p(\mathbf{Y})$ . This models our certainty about the values of the different model variables once the annotations  $\mathbf{Y}$  are observed, and allows us to make predictions on new samples as well as to assess the reliability of the annotators. However, notice that the marginal

$$p(\mathbf{Y}) = \sum_{\mathbf{z}} \int_{\mathbf{f}} \int_{\alpha} \int_{\beta} \int_{\Omega} p(\mathbf{Y}, \mathbf{z}, \mathbf{f}, \alpha, \beta, \Omega) d\mathbf{f} d\alpha d\beta d\Omega \quad (5)$$

190 is not tractable, and therefore we resort to the approximated Variational Bayes (VB) inference procedure.

In principle, inference for this model could be also addressed through Markov Chain Monte Carlo (MCMC) methods, which involve sampling from the pos-

terior instead of approximating it with an explicit probability distribution. In  
 195 fact, MCMC was one of the first approaches for approximate inference in GP  
 [31], and its extension to our model is straightforward from a theoretical view-  
 point. However, MCMC methods are computationally expensive. This issue is  
 exacerbated when using GPs, since the large number of latent variables (at least  
 one for each training instance) and the high correlation that may exist between  
 200 them in the posterior usually requires sophisticated and slow MCMC sampling  
 schemes [32]. Moreover, analytical approximations (such as EP or VB) have  
 obtained excellent results while being significantly faster [33].

In order to approximate the posterior  $p(\mathbf{z}, \Theta | \mathbf{Y})$ , VB minimizes the Kullback-  
 Leibler (KL) divergence with respect to a generic probability distribution  $q(\mathbf{z}, \Theta)$ :

$$\begin{aligned} \text{KL}(q(\mathbf{z}, \Theta) || p(\mathbf{z}, \Theta | \mathbf{Y})) &= \sum_{\mathbf{z}} \int q(\mathbf{z}, \Theta) \log \frac{q(\mathbf{z}, \Theta)}{p(\mathbf{z}, \Theta | \mathbf{Y})} d\Theta \\ &= \sum_{\mathbf{z}} \int q(\mathbf{z}, \Theta) \log \frac{q(\mathbf{z}, \Theta)}{p(\mathbf{Y}, \mathbf{z}, \Theta)} d\Theta + \log p(\mathbf{Y}). \end{aligned}$$

The KL divergence between two distributions is always non negative, and is zero  
 if and only if they coincide. Therefore, the optimal distribution  $q(\mathbf{z}, \Theta)$  in the  
 205 sense of KL divergence minimization is unique and equals the exact  $p(\mathbf{z}, \Theta | \mathbf{Y})$ .  
 Interestingly, notice that we do not need to know the real posterior  $p(\mathbf{z}, \Theta | \mathbf{Y})$   
 to minimize the KL divergence on  $q(\mathbf{z}, \Theta)$ : since  $\log p(\mathbf{Y})$  does not depend on  
 $q(\mathbf{z}, \Theta)$ , only the joint distribution in eq. (4) is required.

However, the sigmoids in  $p(\mathbf{z} | \mathbf{f})$  (recall eq. (1)) prevents us from directly  
 evaluating the KL divergence, since their expectation over a Gaussian cannot be  
 obtained in closed-form. To overcome this problem, a variational lower bound  
 for the sigmoid is used [1, Section 10.6]. Namely, for any  $\xi > 0$ , we have  
 $\sigma(f) = (1 + \exp(-f))^{-1} \geq \sigma(\xi) \exp((f - \xi)/2 - \lambda(\xi)(f^2 - \xi^2))$ , where  $\lambda(\xi) =$   
 $(2\xi)^{-1}(\sigma(\xi) - 1/2)$  [1, Eq. (10.149)]. In our case, this bound yields  $p(\mathbf{z} | \mathbf{f}) \geq$   
 $\mathbf{H}(\mathbf{z}, \mathbf{f}, \boldsymbol{\xi})$ , where

$$\mathbf{H}(\mathbf{z}, \mathbf{f}, \boldsymbol{\xi}) = \prod_{n=1}^N \sigma(\xi_n) \exp \left\{ f_n \left( z_n - \frac{1}{2} \right) - \lambda(\xi_n) f_n^2 + \xi_n^2 \lambda(\xi_n) - \frac{\xi_n}{2} \right\}.$$

Plugging this bound into eq. (4), we have the following lower bound for the joint

distribution

$$p(\mathbf{z}, \Theta, \mathbf{Y}) \geq \mathbf{M}(\mathbf{z}, \mathbf{f}, \alpha, \beta, \Omega, \mathbf{Y}, \xi) = p(\mathbf{Y}|\mathbf{z}, \alpha, \beta) \mathbf{H}(\mathbf{z}, \mathbf{f}, \xi) p(\mathbf{f}|\Omega) p(\alpha) p(\beta) p(\Omega),$$

which in turn produces

$$\text{KL}(q(\mathbf{z}, \Theta) || p(\mathbf{z}, \Theta | \mathbf{Y})) \leq \text{KL}(q(\mathbf{z}, \Theta) || \mathbf{M}(\mathbf{z}, \Theta, \mathbf{Y}, \xi)) + \text{const.} \quad (6)$$

210 Interestingly, notice that  $\mathbf{H}(\mathbf{z}, \mathbf{f}, \xi)$  is quadratic in  $\mathbf{f}$ , which allows us to compute the expectation over a Gaussian in closed-form. Therefore, we focus now on minimizing (with respect to  $q(\mathbf{z}, \Theta)$ ) the analytically tractable right-hand side term in eq. (6), which enforces the left-hand side term (intractable) to be small too. The price for using this bound is a new set of parameters  $\xi = (\xi_1, \dots, \xi_N)^T$   
 215 which need to be estimated.

So far, we have used a generic  $q(\mathbf{z}, \Theta)$  for the approximate posterior distribution. However, VB requires the specification of a particular family, from which the best distribution, in the sense of KL divergence, will be chosen. In this work we use the popular *mean field theory* [1, Section 10.1], which assumes that the approximated distribution factorizes as  $q(\mathbf{z}, \Theta) = q(\mathbf{z})q(\mathbf{f})q(\alpha)q(\beta)q(\Omega)$ . Let  $\bar{\Theta} = \{\mathbf{z}, \Theta\}$  be the set  $\Theta$  expanded with the variable  $\mathbf{z}$ . For  $\theta \in \bar{\Theta}$ , let us write  $\bar{\Theta}_\theta = \bar{\Theta} \setminus \theta$  for the set  $\bar{\Theta}$  minus  $\theta$ , and  $q(\bar{\Theta}_\theta) = \prod_{\eta \in \bar{\Theta}_\theta} q(\eta)$ . Then, for each  $\theta \in \bar{\Theta}$ , it can be shown that the distribution  $q(\theta)$  that minimizes the KL-divergence is given by (see [1, Eq. 10.9] for details)

$$\ln q(\theta) = \langle \ln \mathbf{M}(\mathbf{z}, \Theta, \mathbf{Y}, \xi) \rangle_{q(\bar{\Theta}_\theta)} + \text{const.} \quad (7)$$

Alternating the estimation of  $q(\mathbf{z})$ ,  $q(\mathbf{f})$ ,  $q(\alpha)$ ,  $q(\beta)$  and  $q(\Omega)$  leads to an iterative algorithm where the KL divergence decreases after each iteration. Since it is always a non-negative number, the convergence is ensured.

To calculate  $q(\mathbf{z})$ , we deduce from eq. (7) that it factorizes as  $q(\mathbf{z}) = \prod_{n=1}^N q(z_n)$ . Thus, we can compute each  $q(z_n)$  separately. Since  $z_n$  only takes

two values, we have

$$\begin{aligned} q(z_n = 0) &\propto \prod_{r \in R_n} \exp \{y_n^r \langle \log(1 - \beta_r) \rangle + (1 - y_n^r) \langle \log \beta_r \rangle\}, \\ q(z_n = 1) &\propto \exp(\langle f_n \rangle) \prod_{r \in R_n} \exp \{y_n^r \langle \log \alpha_r \rangle + (1 - y_n^r) \langle \log(1 - \alpha_r) \rangle\}. \end{aligned} \quad (8)$$

For  $q(\mathbf{f})$  we observe that  $\langle \ln \mathbf{M}(\mathbf{z}, \Theta, \mathbf{Y}, \xi) \rangle_{q(\Theta_f)}$  cannot be calculated. To avoid this problem, we assume that  $q(\Omega)$  is a degenerate distribution. Then,  $\langle \ln \mathbf{M}(\mathbf{z}, \Theta, \mathbf{Y}, \xi) \rangle_{q(\Theta_f)}$  becomes a quadratic function of  $\mathbf{f}$  and, therefore, the approximate posterior  $q(\mathbf{f})$  is a Gaussian  $\mathcal{N}(\mathbf{f} | \boldsymbol{\mu}_f, \boldsymbol{\Sigma}_f)$ . Mean and covariance are calculated by taking first and second order derivatives of  $\log q(\mathbf{f})$  to obtain:

$$\boldsymbol{\mu}_f = \boldsymbol{\Sigma}_f (\langle \mathbf{z} \rangle - (\mathbf{1}/2)), \quad \boldsymbol{\Sigma}_f = (\mathbf{K}^{-1} + 2\boldsymbol{\Lambda})^{-1}, \quad (9)$$

where  $\boldsymbol{\Lambda} = \text{diag}(\lambda(\xi_1), \dots, \lambda(\xi_N))$ , and  $\langle \mathbf{z} \rangle = (q(z_1 = 1), \dots, q(z_n = 1))^T$ .

Since  $q(\Omega)$  is a degenerate distribution, we only need the value of  $\Omega$  where  $q(\Omega)$  is not zero. For that, we minimize the following objective function

$$\mathcal{L}(\Omega) = \ln |\mathbf{K} + (2\boldsymbol{\Lambda})^{-1}| + \mathbf{u}^T (\mathbf{K} + (2\boldsymbol{\Lambda})^{-1})^{-1} \mathbf{u}, \quad (10)$$

220 where  $\mathbf{u} = (\mathbf{1}/2) \cdot \boldsymbol{\Lambda}^{-1} (\langle \mathbf{z} \rangle - (\mathbf{1}/2))$ . Recall also that  $\mathbf{K}$  depends on  $\Omega$ .

To calculate  $q(\boldsymbol{\alpha})$  and  $q(\boldsymbol{\beta})$ , we deduce from eq. (7) that both factorize as  $q(\boldsymbol{\alpha}) = \prod_{r=1}^R q(\alpha_r)$  and  $q(\boldsymbol{\beta}) = \prod_{r=1}^R q(\beta_r)$ . Then we can calculate each  $q(\alpha_r)$  and  $q(\beta_r)$  separately. From eq. (7) we obtain the following Beta distributions:

$$q(\alpha_r) = \text{Beta} \left[ \alpha_r \left| a_0^\alpha + \sum_{n \in N_r} \langle z_n \rangle y_n^r, b_0^\alpha + \sum_{n \in N_r} \langle z_n \rangle (1 - y_n^r) \right. \right], \quad (11)$$

$$q(\beta_r) = \text{Beta} \left[ \beta_r \left| a_0^\beta + \sum_{n \in N_r} (1 - \langle z_n \rangle) (1 - y_n^r), b_0^\beta + \sum_{n \in N_r} (1 - \langle z_n \rangle) y_n^r \right. \right]. \quad (12)$$

For  $\xi$  we maximize  $\langle \ln \mathbf{M}(\Theta, \mathbf{Y}, \mathbf{X}, \xi) \rangle_{q(\Theta)}$  w.r.t. each  $\xi_n$ , which yields

$$\xi_n = \sqrt{\langle f_n \rangle^2 + \boldsymbol{\Sigma}_f(n, n)}. \quad (13)$$

The whole estimation procedure is summarized in Algorithm 1. Notice that an initial approximated posterior for the true labels  $q^0(\mathbf{z})$  is required. We propose to initialize it with *soft majority voting*, that is,  $q^0(z_n = 1)$  is the proportion

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**Algorithm 1** VGPCR (Variational GP for CRowdsourcing)

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**Require:**  $\mathbf{X}, \mathbf{Y}, \boldsymbol{\xi}^0 = \mathbf{1}, q^0(\mathbf{z}), k = 0.$

- 1: **repeat**
  - 2:   Calculate  $\boldsymbol{\Omega}^{k+1}$  as the minimizer of eq. (10) using  $q^k(\mathbf{z})$  and  $\boldsymbol{\xi}^k$ ;
  - 3:   Update  $q^{k+1}(\mathbf{f})$  with eq. (9) using  $q^k(\mathbf{z}), \boldsymbol{\xi}^k$ , and  $\boldsymbol{\Omega}^{k+1}$ ;
  - 4:   Update  $q^{k+1}(\boldsymbol{\alpha})$  and  $q^{k+1}(\boldsymbol{\beta})$  with eqs. (11)-(12) using  $q^k(\mathbf{z})$  ;
  - 5:   Update  $q^{k+1}(\mathbf{z})$  with eq. (8) using  $q^{k+1}(\mathbf{f})$ . The expectations are calculated with eq. (3) using  $q^{k+1}(\boldsymbol{\alpha})$  and  $q^{k+1}(\boldsymbol{\beta})$ .
  - 6:   Calculate  $\boldsymbol{\xi}^{k+1}$  with eq. (13) using  $q^{k+1}(\mathbf{f})$ ;
  - 7:    $k = k + 1$ ;
  - 8: **until** convergence
  - 9: **output**  $q(\mathbf{z}, \Theta)$
- 

of annotators that assign label 1 to the sample  $\mathbf{x}_n$ . This initialization implicitly  
225 assumes that most of the annotators are not adversarial. Otherwise, due to the  
ambiguity of eq. (2), we would train a classifier predicting the opposite labels.

#### 4. The predictive distribution

Once the model is trained, we are given a new sample  $\mathbf{x}_*$  and we need to predict the probability of each class. In a crowdsourcing problem, we additionally  
230 might have access to a set of labels  $\mathbf{y}_* = \{y_*^r \in \{0, 1\} : r \in R_* \subseteq \{1, \dots, R\}\}$   
provided by some annotators. To the best of our knowledge, this plausible  
scenario has not been addressed before in the crowdsourcing literature.

In our model, the prediction can be naturally obtained as the conditional of the hidden label  $z_*$  given the observed labels  $\mathbf{y}_*$  and  $\mathbf{Y}$ , that is,

$$p(z_* | \mathbf{y}_*, \mathbf{Y}) = \sum_{\mathbf{z}} \int p(z_*, \mathbf{z}, f_*, \mathbf{f}, \boldsymbol{\alpha}, \boldsymbol{\beta}, \boldsymbol{\Omega} | \mathbf{y}_*, \mathbf{Y}) d\boldsymbol{\Theta} df_* \approx \tag{14}$$
$$\text{const} \cdot \left[ \int p(z_* | f_*) \left( \int p(f_* | \mathbf{f}) q(\mathbf{f}) d\mathbf{f} \right) df_* \right] \cdot \left[ \int p(\mathbf{y}_* | z_*, \boldsymbol{\alpha}, \boldsymbol{\beta}) q(\boldsymbol{\alpha}) q(\boldsymbol{\beta}) d\boldsymbol{\alpha} d\boldsymbol{\beta} \right].$$

The GP conditional is  $p(f_* | \mathbf{f}) = \mathcal{N}(f_* | \mathbf{h}^T \mathbf{K}^{-1} \mathbf{f}, c - \mathbf{h}^T \mathbf{K}^{-1} \mathbf{h})$ , where  $\mathbf{h} = [k(\mathbf{x}_1, \mathbf{x}_*), k(\mathbf{x}_2, \mathbf{x}_*), \dots, k(\mathbf{x}_N, \mathbf{x}_*)]^T$ ,  $c = k(\mathbf{x}_*, \mathbf{x}_*)$ , and  $\mathbf{K}$  is the kernel matrix

235 in  $\mathbf{X}$ . Then, using  $q(\mathbf{f}) = \mathcal{N}(\mathbf{f}|\boldsymbol{\mu}_f, \boldsymbol{\Sigma}_f)$  (recall eq. (9)) we obtain  $\int p(f_*|\mathbf{f})q(\mathbf{f})d\mathbf{f} = \mathcal{N}(f_*|m_*, s_*^2)$ , where  $m_* = \mathbf{h}^T \mathbf{K}^{-1} \boldsymbol{\mu}_f$  and  $s_*^2 = c - \mathbf{h}^T (\mathbf{K} + (2\boldsymbol{\Lambda})^{-1})^{-1} \mathbf{h}$ .

Substituting back in eq. (14) and using eq. (4.153) in [1], we obtain the following predictive distribution for  $z_*$ :

$$p(z_* = 1|\mathbf{y}_*, \mathbf{Y}) \propto \sigma(\kappa(s_*^2)m_*) \prod_{r \in R_*} \langle \alpha_r \rangle_{q(\alpha_r)}^{y_*^r} (1 - \langle \alpha_r \rangle_{q(\alpha_r)})^{1-y_*^r}, \quad (15)$$

$$p(z_* = 0|\mathbf{y}_*, \mathbf{Y}) \propto (1 - \sigma(\kappa(s_*^2)m_*)) \prod_{r \in R_*} (1 - \langle \beta_r \rangle_{q(\beta_r)})^{y_*^r} \langle \beta_r \rangle_{q(\beta_r)}^{1-y_*^r}, \quad (16)$$

where  $\kappa(s_*^2) = (1 + \pi s_*^2/8)^{-1/2}$ . Notice that this distribution generalizes the case where no information is provided by the annotators, that is  $\mathbf{y}_* = \emptyset$ . In such a case, the predictive distribution for  $z_*$  is the Bernoulli distribution  $p(z_*|\mathbf{Y}) = [\sigma(\kappa(s_*^2)m_*)]^{z_*} [1 - \sigma(\kappa(s_*^2)m_*)]^{1-z_*}$ . Finally, a threshold  $\delta$  is used on  $p(z_* = 1|\mathbf{y}_*, \mathbf{Y})$  to assign the new sample  $\mathbf{x}_*$  to  $\mathcal{C}_1$ .

In the next section we will compare our novel crowdsourcing method against current state-of-the-art approaches. We will observe that the proposed method stands out as the most robust approach across a wide range of datasets. In particular, we will see that the proposed VB inference is better suited than EP  
 245 for GP-based crowdsourcing classifiers. Finally, although the main goal is to illustrate the performance of the proposed method, we will also examine the behavior, strengths, and weaknesses of the other methods it is compared with. To some extent, this provides an up-to-date experimental review of the main  
 250 crowdsourcing approaches in the literature.

## 5. Experimental Results

In this section, we provide a comprehensive experimentation that compares the proposed method with several state-of-the-art approaches on three different types of datasets. First, we make use of *fully synthetic* data, where crowdsourc-  
 255 ing annotations are synthetically generated for an also synthetic underlying classification dataset. This constitutes a completely controlled framework where we can check the expected behavior of the compared algorithms. Second, we evaluate the methods on *semi-synthetic* data, where the underlying classification

Table 2: The three types of data used in this work.

	<b>Fully synthetic</b>	<b>Semi-synthetic</b>	<b>Fully real</b>
<i>Classif. data set</i>	Synthetic	Real	Real
<i>Annotations</i>	Synthetic	Synthetic	Real
<i>Examples</i>	1D cosine-based	Heart Sonar	Sentence Polarity Music Genre

dataset comes from a real-world problem but the crowdsourcing annotations  
 260 are obtained synthetically. This is an interesting and popular hybrid setting in  
 crowdsourcing, where we can keep the influence of the real underlying classifi-  
 cation dataset apart from the crowdsourced annotations, which remain under  
 control. Third, we evaluate the methods on *fully real* data, where both features  
 and annotations come from a real problem. This is the most realistic setting for  
 265 practical applications, although we have no knowledge about the data genera-  
 tion process. Table 2 summarizes the types of data used in this work.

The proposed method is referred to as *VGPCR* (Variational GP for CRowd-  
 sourcing). In all the experiments, *VGPCR* is compared against the state-of-the-  
 art GP-based crowdsourcing method in [12] (*Rodrigues*), which utilizes EP as  
 270 inference procedure. In the comparison we also include the most straightforward  
 manner to apply a GP to the crowdsourcing setting, *GP-MV*, which consists of a  
 standard GP classifier trained with the Majority Voting (MV) labels. The last  
 GP-based method included in the comparison is a GP classifier trained with  
 the true labels (*GP-GOLD*)<sup>2</sup>. Notice that, intuitively, *GP-GOLD* and *GP-MV*  
 275 provide (respectively) upper and lower bounds for the GP-based crowdsourcing  
 methods *Rodrigues* and *VGPCR*. Finally, to obtain a more thorough compari-  
 son, we include the methods in [9] (*Raykar*), and [10] (*Yan*), which are based

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<sup>2</sup>Clearly, *GP-GOLD* can only be trained if there are real labels available for the training  
 set. Of course, this is not common in a real crowdsourcing application (otherwise it could be  
 cast as a standard classification problem). However, in the two real datasets used here the  
 true labels are also provided in order to compare with *GP-GOLD*.

in LR instead of GP (recall the third paragraph in Section 1).

If the annotators provide labels for the test set (that is, some  $\mathbf{y}_*$  are available), then our method is referred to as *VGPCR\**. As a baseline, we find interesting to compare *VGPCR\** with the most straightforward way to predict with the test set annotations, which we refer to as *MV\**, and whose predictions are based only on these annotations (no training step is needed). A brief summary of all the algorithms used in the experiments is provided in Table 3.

Table 3: An overview of the methods compared in the experiments. From top to bottom, the thick horizontal lines separate non-crowdsourcing methods (*GP-GOLD*), crowdsourcing algorithms that do not use test set annotations, and approaches that do use them.

Algorithm	Description
<i>GP-GOLD</i>	Intuitive upper bound for the GP-based crowdsourcing methods. Trains a GP with the real labels (it is not a crowdsourcing algorithm).
<i>GP-MV</i>	Simplest way to apply GP to crowdsourcing (intuitive lower bound). Trains a GP with the majority voting labels.
<i>Rodrigues</i>	State-of-the-art GP-based crowdsourcing method proposed in [12]. EP inference is used.
<i>VGPCR</i>	GP-based crowdsourcing method <i>proposed here</i> . Variational inference is used.
<i>Raykar</i>	Based on logistic regression. EM for inference. Proposed in [9]. First probabilistic model for crowdsourcing.
<i>Yan</i>	Based on logistic regression. EM for inference. Proposed in [10]. Annotators parameters depend on the instance they label.
<i>MV*</i>	Simplest (naive) way to use test set annotations for prediction. It does not need a training step.
<i>VGPCR*</i>	Straightforward extension to <i>VGPCR</i> . <i>Proposed here</i> . Probabilistically integrates test set annotations in the prediction.

The predictive performance of the methods is compared using two popular metrics: the area under the ROC curve (AUC), and the overall accuracy (OA), which is calculated for the threshold  $\delta = 1/2$ . Moreover, in order to compare the computational cost, the CPU time needed to train each method is also provided.

We implemented  $VGPCR^{(*)}$ , *Raykar*, *Yan*, and GP-classification (necessary  
 290 for *GP-GOLD* and *GP-MV*) in Matlab<sup>©</sup>, whereas a Matlab<sup>©</sup> implementation  
 for *Rodrigues* can be downloaded from his website <http://www.fprodrigues.com>. All the code and datasets are available at <http://decsai.ugr.es/vip/software.html>. The experiments were run on the same machine Intel<sup>©</sup> Xeon<sup>©</sup>  
 E5-4640 @ 2.40GHz.

### 295 5.1. Fully synthetic data

In this section we compare the performance of the methods with a controlled  
 one-dimensional example. Figure 2a) shows the underlying synthetic classifica-  
 tion dataset used. The features are uniformly sampled in the interval  $[-\pi, \pi]$ .  
 The real labels are assigned according to the sign of the cosine function on each  
 300 sample: class  $\mathcal{C}_1$  (resp. class  $\mathcal{C}_0$ ) if the cosine is positive (resp. negative). Then,  
 we simulate  $R = 5$  annotators by fixing the values of sensitivity and specificity  
 to  $\boldsymbol{\alpha} = \{0.9, 0.7, 0.8, 0.1, 0.9\}$  and  $\boldsymbol{\beta} = \{0.6, 0.8, 0.5, 0.2, 0.8\}$ , respectively. That  
 is, if the true label of the  $n$ -th sample is  $z_n = 1$  (resp.  $z_n = 0$ ), the  $r$ -th annota-  
 tor assigns it to class  $\mathcal{C}_1$  (resp.  $\mathcal{C}_0$ ) with probability  $\alpha_r$  (resp.  $\beta_r$ ). In Fig. 2 (b-f)  
 305 we show the labels assigned by each annotator. As expected from the values of  
 $\boldsymbol{\alpha}$  and  $\boldsymbol{\beta}$ , annotators 1, 2, 3, and 5 make fewer mistakes than annotator 4, who  
 assigns most samples to the opposite class (it has an *adversarial* behavior).

The experiment is repeated 10 times with different training sets of 100 sam-  
 ples (50 of each class). In each realization we also generate a uniformly sampled  
 310 test set with 200 instances (100 each class). Moreover, test set annotations are  
 also simulated in order to apply the  $MV^*$  and  $VGPCR^*$  algorithms.

Table 4 shows the predictive performance of all the methods for the 10  
 realizations. Let us focus first on the five crowdsourcing algorithms that do not  
 use the test set annotations (i.e., the central columns of the table). The results  
 315 show two clear groups: those based on GP (*GP-MV*, *Rodrigues*, *VGPCR*), whose  
 results are competitive with *GP-GOLD*, and those that use LR (*Raykar*, *Yan*),  
 whose performance is really poor. This is a reasonable behavior if we take into  
 account that LR decision boundaries are hyperplanes, i.e., one point in this 1-D

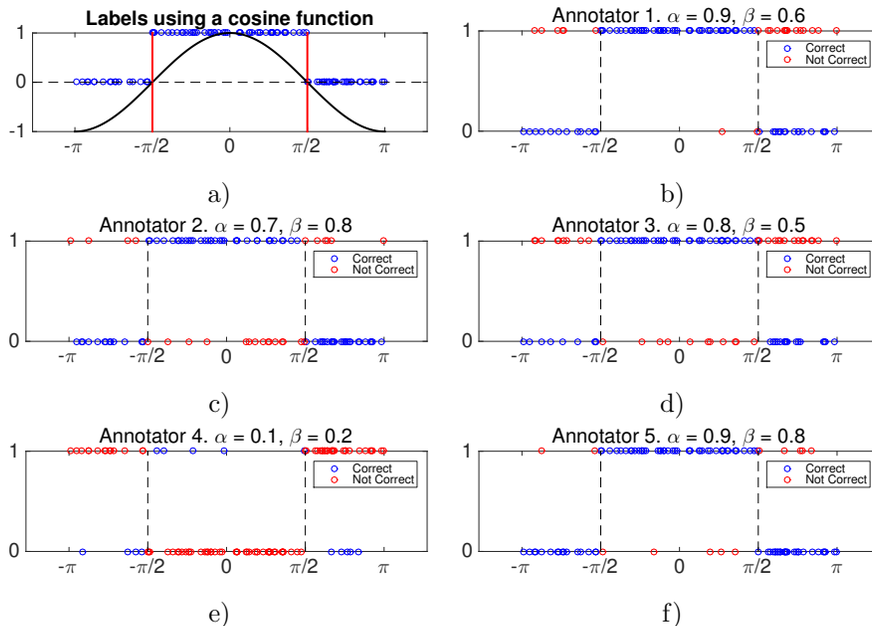


Figure 2: a) Original data set labeled using sign of cosine function. b) - f) Labels provided by annotators 1,2,3,4 and 5 respectively.

example. This is clearly insufficient to deal with our training dataset, where  $\mathcal{C}_0$  has two disconnected parts with  $\mathcal{C}_1$  in the middle (recall fig. 2a).

Among the LR-based algorithms, we observe that *Yan* performs considerably better than *Raykar*. This means that *Yan*'s feature-dependent model for the annotations is, to some extent, helping to compensate for the insufficient LR model. It is worth noting that the mean result for *Raykar* is hardly above a random guess (around 0.5 of AUC and OA). This LR deficiency is clearly overcome by the GP-based methods, which manage to effectively separate the classes by using a non-linear kernel that allows for more complex decision boundaries (SE kernel in this work, recall Section 2). Among the GP-based methods, the proposed *VGPCR* obtains the best result, followed closely by *Rodrigues*. Notice that *GP-MV* is very close to them in AUC but not in OA, which implies that the threshold  $\delta = 1/2$  is not the most appropriate one for class prediction in *GP-MV* (although the classes are well-separated by some other threshold). Nonetheless, this simple 1-D example turns out to be too easy for the GP-based

Table 4: Predictive performance of the compared methods for the 10 realizations of the fully synthetic experiment. The best mean performance among the crowdsourcing methods that do not use test set annotations (central columns) is bolded.

Rea.	<i>GP-GOLD</i>		<i>GP-MV</i>		<i>Rodrigues</i>		<i>VGPCR</i>		<i>Raykar</i>		<i>Yan</i>		<i>MV*</i>		<i>VGPCR*</i>	
	AUC	OA%	AUC	OA%	AUC	OA%	AUC	OA%	AUC	OA%	AUC	OA%	AUC	OA%	AUC	OA%
1	1.0000	100.00	0.9978	68.00	1.0000	100.00	1.0000	99.50	0.5800	50.00	0.5800	79.00	0.8520	77.00	1.0000	100.00
2	1.0000	98.50	1.0000	84.50	1.0000	98.50	1.0000	98.50	0.4400	50.00	0.5600	57.00	0.8572	78.00	0.9998	99.00
3	1.0000	100.00	1.0000	64.50	1.0000	100	1.0000	99.50	0.4200	50.00	0.7695	75.50	0.7836	76.00	1.0000	100.00
4	0.9998	98.50	0.9987	94.50	1.0000	94.50	1.0000	96.50	0.5000	50.00	0.5000	50.00	0.8357	75.50	1.0000	100.00
5	1.0000	99.00	0.9934	90.50	0.9965	94.50	0.9984	97.00	0.4700	50.00	0.5654	71.00	0.8435	79.00	1.0000	99.50
6	1.0000	100.00	0.9971	91.00	1.0000	100	1.0000	100.00	0.5100	69.00	0.7403	75.00	0.8261	77.00	1.0000	99.00
7	1.0000	99.00	0.9974	96.00	0.9993	98.00	0.9996	98.00	0.5700	76.00	0.6267	71.50	0.8171	76.50	1.0000	99.50
8	0.9990	97.50	0.9928	94.00	0.9972	95.50	0.9983	97.50	0.5500	46.50	0.7435	75.00	0.8436	77.00	0.9997	98.50
9	1.0000	99.50	0.9895	91.00	0.9997	98.00	1.0000	99.50	0.5100	41.50	0.5100	68.00	0.8368	78.50	1.0000	99.50
10	1.0000	98.00	0.9970	78.50	0.9999	98.00	0.9999	96.50	0.4900	50.00	0.7450	73.00	0.8419	79.50	1.0000	99.50
Mean	0.9999	99.00	0.9964	85.25	0.9993	97.70	<b>0.9996</b>	<b>98.25</b>	0.5040	53.30	0.6340	69.50	0.8337	77.40	1.0000	99.45

methods, and further differences will be appreciated in subsequent experiments.

335 It is also interesting to check that, as theoretically intuited, *Rodrigues* and *VGPCR* performances are upper and lower bounded by *GP-GOLD* and *GP-MV*, respectively. Moreover, the differences with *GP-GOLD* are almost insignificant, which means that the crowdsourcing methods are able to extract from the noisy annotations almost the same information as a full GP does from the true labels.

340 Let us now concentrate on the methods that use test set annotations (*MV\** and *VGPCR\**). The latter reaches mean AUC and OA of 1.0000 and 99.45% respectively, and manages to totally separate the classes in 8 out of the 10 realizations. These results are better than those obtained by *VGPCR*, which supports the idea that crowdsourcing methods can benefit from the probabilistic  
345 integration of test set annotations if available. Moreover, notice that *VGPCR\** even outperforms *GP-GOLD* (and this will be also the case in subsequent experiments). In fact, this is expected when the annotations generation process follows the one proposed in the model (like in the synthetic and semi-synthetic experiments) and the sensitivity and specificity parameters ( $\alpha$ ,  $\beta$ ) are correctly  
350 estimated in the training step (as we will check in the following paragraph). In that case, test annotations are a very valuable source of information for *VG-PCR\**, as they directly depend on the true test label through  $\alpha$  and  $\beta$ . This has an interesting practical implication in real problems: as long as the crowdsour-

Table 5: Estimated values of sensitivity and specificity for the five annotators in the fully synthetic experiment. Only those methods that include these parameters in their formulation are shown. The values are the mean over the 10 realizations.

Annotator	Original		<i>Raykar</i>		<i>Rodrigues</i>		<i>VGPCR</i>	
	$\alpha$	$\beta$	$\alpha$	$\beta$	$\alpha$	$\beta$	$\alpha$	$\beta$
1	0.9	0.6	0.8910	0.6340	0.8183	0.5433	0.8993	0.6128
2	0.7	0.8	0.6855	0.8410	0.6058	0.7461	0.6828	0.8119
3	0.8	0.5	0.7916	0.4880	0.7607	0.4487	0.8019	0.4837
4	0.1	0.2	0.1362	0.1793	0.2081	0.2724	0.1015	0.1823
5	0.9	0.8	0.8908	0.8168	0.7986	0.6920	0.9042	0.7816

ing annotation model is representative for the data at hand, it is more useful to  
 355 collect non-experts opinions for test instances than to collect expert true labels  
 for train ones. Regarding  $MV^*$ , its performance is clearly below  $VGPCR^*$ . This  
 is reasonable since  $MV^*$  does not consider any probabilistic model for the anno-  
 tations, and thus it is very sensitive to the presence of noise in them (indeed, it  
 performs better as  $\alpha, \beta \rightarrow 1$ , i.e., when the noise decreases and the annotations  
 360 themselves become very representative of the underlying true labels).

Finally, Table 5 shows the estimated values of sensitivity and specificity for  
 the models that include them in their formulation (i.e., *Rodrigues*, *VGPCR*,  
 and *Raykar*, since *Yan* uses a more complex feature-dependent model). The  
 proposed *VGPCR* method obtains the most accurate estimations: a maximum  
 365 absolute difference of 0.0184, whereas it is 0.0410 for *Raykar* and 0.1081 for  
*Rodrigues* (in next Section 5.2 we will analyze the difficulties of *Rodrigues* to es-  
 timate  $\alpha$  and  $\beta$ ). As stated in the previous paragraph, these reliable estimations  
 of  $\alpha/\beta$  in *VGPCR* imply that  $VGPCR^*$  greatly benefits from test annotations.  
 Moreover, the estimations of *VGPCR* for annotator 4 are quite accurate, which  
 370 means that it has been able to recognize its adversarial behavior (*Rodrigues* and  
*Raykar* also detect it, although less accurately, especially *Rodrigues*). Finally,  
 we stress that the poor performance reported for *Raykar* in Table 4 does not  
 come from a wrong estimation of  $\alpha$  or  $\beta$ , but from the underlying LR modeling.

For this simple synthetic experiment, CPU training time is not reported,  
375 since all values are very similar (there are only 100 training 1-D instances).

## 5.2. Semi-synthetic data

In this section we follow an analogous experimental approach as before, but  
focusing on two more complex semi-synthetic datasets. This allows us to gain  
additional insight into the behavior of the compared methods. In particular, we  
380 observe that the proposed method *VGPCR* (and *VGPCR\**) stand out as the  
most effective and robust crowdsourcing approaches across the two experiments.

### 5.2.1. Heart dataset

This database, also known as *Heart Disease*, is a popular real classifica-  
tion problem donated by the Cleveland Clinic Foundation to the UCI Machine  
385 Learning repository, see [http://archive.ics.uci.edu/ml/datasets/Heart+](http://archive.ics.uci.edu/ml/datasets/Heart+Disease)  
[Disease](http://archive.ics.uci.edu/ml/datasets/Heart+Disease). The goal is to predict the presence or absence (i.e., binary problem) of  
heart disease in the patient. For that, it contains 13 relevant explanatory vari-  
ables (features), such as age, resting blood pressure, and maximum heart rate.  
After discarding 6 instances with missing features, the final dataset contains  
390 297 samples (137 with disease and 160 without it).

With this real underlying classification problem, we simulate  $R = 5$  crowd-  
sourcing annotators with the same sensitivity and specificity values as before,  
i.e.,  $\alpha = \{0.9, 0.7, 0.8, 0.1, 0.9\}$  and  $\beta = \{0.6, 0.8, 0.5, 0.2, 0.8\}$ . Notice that  
the crowdsourcing setting is very appropriate for this medical domain, where  
395 different doctors (annotators) may have different opinions (annotations) about  
the presence/absence of heart disease based on the 13 provided features. The  
adversarial behavior of annotator 4 represents the meddling of a non-expert an-  
notator who is confusing both classes. We will see that crowdsourcing methods  
are able to identify this type of undesirable annotator, and take advantage of  
400 their opinions in light of their degree of expertise.

To average the results over different runs, we consider 10 independent ran-  
dom train/test partitions with 208/89 instances respectively. The results are

shown in Table 6 and Figure 3. The table contains the AUC and OA mean values for both the test and train datasets. Moreover, it shows the mean CPU time needed to train each method. In the figure we focus on two of these quantities, analyzing the trade-off between generalization capability (in terms of test AUC) and computational cost (in terms of CPU train time). The figure does not include the methods that use test set annotations. Moreover, the figure displays plus/minus one standard deviation of the shown mean quantities. Finally, Table 7 presents the estimated specificity and sensitivity values.

Table 6: Results in the *heart* semi-synthetic dataset. Test and train performances (in terms of AUC and OA) and the CPU time needed to train each method are provided. The results are the mean over the 10 runs. The best generalization (test) performance among the crowdsourcing methods that do not use test set annotations (central rows) is bolded.

Methods	Test set		Train set		CPU time (s)
	AUC	OA%	AUC	OA%	
<i>GP-GOLD</i>	0.8898	81.91	0.9349	86.25	120.65
<i>GP-MV</i>	0.8633	69.33	0.9133	75.91	46.36
<i>Rodrigues</i>	0.8239	78.09	0.9827	93.46	913.06
<i>VGPCR</i>	<b>0.8870</b>	<b>82.02</b>	0.9298	86.20	29.21
<i>Raykar</i>	0.8853	80.34	0.9287	86.01	0.54
<i>Yan</i>	0.7396	63.37	0.7944	72.69	625.06
<i>MV*</i>	0.8211	74.04	0.8350	76.88	0
<i>VGPCR*</i>	0.9921	95.62	0.9935	96.30	29.21

We observe that, among the five crowdsourcing methods, the proposed *VG-PCR* gets the best generalization (test) performance in both AUC and OA. In fact, in the latter metric it even outperforms *GP-GOLD*, which is trained with the true labels. This means that our method is making the most of the noisy labels that is provided with, reaching the level of its intuitive upper bound. Table 7 also supports that *VGPCR* is able to accurately figure out the model that generates the annotations. The estimated values for  $\alpha$  and  $\beta$  are very close to the true original ones (better than those obtained by *Rodrigues* and similar

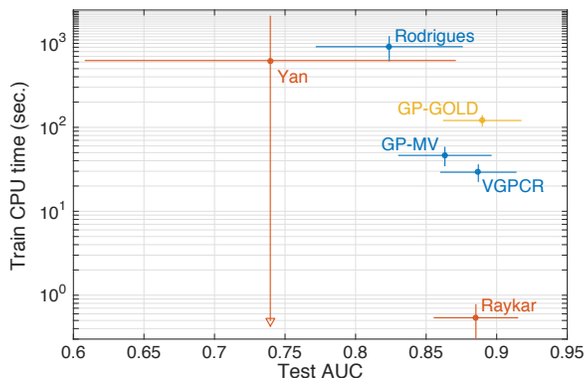


Figure 3: Trade-off between generalization capability (in terms of test AUC) and computational cost (in terms of CPU train time) in the *heart* dataset. For each method, the mean plus/minus one standard deviation is shown. The color indicates the family of the algorithm, i.e. yellow for the non-crowdsourcing method *GP-GOLD*, blue for the GP-based crowdsourcing methods, and red for the LR-based ones. Notice also the logarithmic scale in the y-axis.

to *Raykar*'s). In particular, it manages to detect the adversarial behavior of  
 420 annotator 4. Moreover, it obtains the second shortest CPU train time, 29.21 s.

Regarding its GP-based competitors, the intuitive lower bound *GP-MV* exhibits a worse predictive capacity, as expected. The behavior of *Rodrigues* is, however, more surprising and worth analyzing. *Rodrigues* obtains quite poor test performance, far from *GP-GOLD*, *VGPCR*, and even its supposed lower  
 425 bound *GP-MV*. The key is given by its performance in the training set. There, we observe that *Rodrigues* is fitting very well the training instances (e.g. 0.9827 of train AUC), much better than the rest of algorithms (even *GP-GOLD*). This is the so-called *over-fitting* problem, which happens when a machine learning  
 430 method fits the training data too faithfully, at the expense of its generalization capability. It is also worth pointing that *Rodrigues* is the most computationally heavy method (and with a reduced standard deviation, see Figure 3). We will see that this inefficiency of the EP inference is recurrent across all experiments, and in Section 5.3.1 we will analyze it in more detail. In the comparison with *VGPCR*, it is also interesting to note that the  $\alpha$  and  $\beta$  estimates provided by

Table 7: Estimated values of sensitivity and specificity for the five annotators in the *heart* semi-synthetic experiment. Only those methods that include these parameters in their formulation are shown. The values are the mean over the 10 realizations.

Annotator	Original		<i>Raykar</i>		<i>Rodrigues</i>		<i>VGPCR</i>	
	$\alpha$	$\beta$	$\alpha$	$\beta$	$\alpha$	$\beta$	$\alpha$	$\beta$
1	0.9	0.6	0.8862	0.6070	0.7058	0.4417	0.8901	0.6065
2	0.7	0.8	0.7052	0.7995	0.5144	0.6310	0.7123	0.8021
3	0.8	0.5	0.8151	0.4772	0.7151	0.3821	0.8177	0.4772
4	0.1	0.2	0.1065	0.1789	0.3808	0.4259	0.1006	0.1790
5	0.9	0.8	0.8807	0.7834	0.6271	0.5554	0.8904	0.7877

435 *Rodrigues* are clearly less accurate than those obtained by the proposed method, see Table 7. From a practical viewpoint, this means that *Rodrigues* faces difficulties to identify the annotators reliability, which is a very relevant information for the user. We hypothesize that all this enhancement is due to the two main differences between *Rodrigues* and *VGPCR*: the use of variational inference and  
440 the more refined modeling of the annotators. In subsequent experiments, we will further support this idea and analyze some other subtle differences in the modeling of *VGPCR* and *Rodrigues*.

Next, let us analyze the results of the LR-based methods. Interestingly, the simpler model of *Raykar* obtains an excellent test performance, very close to  
445 the GP-based *VGPCR* and even *GP-GOLD*. This suggests some latent linear structure in the *heart* dataset. Otherwise, as we saw in the fully synthetic experiment, the LR hyperplanes could not produce as good results as the more complex GP boundaries. In order to confirm this linearity in the underlying *heart* dataset, we trained a standard LR classifier with the true labels (following the same scheme as for the GP-based *GP-GOLD*). As expected, the mean  
450 test AUC is 0.8884, almost the same as *GP-GOLD* (recall Table 6). Moreover, *Raykar*'s estimations for  $\alpha$  and  $\beta$  are quite accurate (Table 7), and its computational cost is insignificant (it is the fastest method). Therefore, it could be stated that, in this close-to-linear dataset, the LR-based *Raykar* gets the

455 best trade-off between predictive performance and computational cost. Even  
better than the proposed GP-based *VGPCR* (see Figure 3). This supports the  
common practice in Machine Learning that, when data is simple, a good model  
does not need to be a complicated one. However, in the remaining experiments  
we will find more complex datasets where *Raykar* cannot keep up with the level  
460 of the proposed *VGPCR*.

On the other hand, in spite of the aforementioned linearity, *Yan* obtains a  
very poor (the worst) test performance in this dataset. This must be a conse-  
quence of its more complex feature-dependent model for the annotations, which  
makes the convergence at the training step more challenging. This is reflected  
465 in the large standard deviations exhibited by *Yan* in Figure 3, which show that  
different runs have converged to very different parameters, leading to very het-  
erogeneous results<sup>3</sup>. Moreover, recall that the synthetic generation process used  
for the annotations does not depend on the features. Therefore, this scenario  
seems more favorable to *Raykar*, and it will be convenient to compare both  
470 methods in the fully real datasets.

Regarding the methods that use test set annotations, the conclusions are  
the same as in the previous section. Again, *VGPCR\** obtains an almost perfect  
separation between classes, which is mainly caused by the accurate estimation  
of  $\alpha$  and  $\beta$  (recall Table 7). We also observe that the baseline *MV\** is not  
475 competitive against *VGPCR\**, as it is very sensitive to the noisy labels. Recall  
that *MV\** does not need a training step (therefore, its CPU train time is 0).

### 5.2.2. Sonar dataset

This database, also known as *Sonar, Mines vs Rocks*, is a real classifica-  
tion problem donated by R.P. Gorman and T.J. Sejnowski to the UCI Ma-  
chine Learning repository, see <http://archive.ics.uci.edu/ml/datasets/>  
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<sup>3</sup>In particular, we observed that the surprisingly high mean training CPU time for *Yan* is  
mainly caused by 2 of the 10 runs, which really struggled to converge. Without them, the  
mean would be 26.14 seconds, more in accordance with the other LR-based *Raykar*.

`connectionist+bench+(sonar,+mines+vs.+rocks)`. The goal is to distinguish between rocks and mines (metal cylinders) by analyzing the sonar signals bounced off these materials. The features are 60 numbers in the range  $[0, 1]$ , where each number represents the energy within a particular frequency band. The dataset includes 208 records, 97 samples correspond to rocks and 111 to mines. For this real underlying classification problem, we simulate  $R = 5$  crowdsourcing annotators with the same sensitivity and specificity values as before, i.e.,  $\alpha = \{0.9, 0.7, 0.8, 0.1, 0.9\}$  and  $\beta = \{0.6, 0.8, 0.5, 0.2, 0.8\}$ .

To average the results over different runs, we consider 10 independent random train/test partitions with 146/62 instances. As in the *heart* dataset, the results are shown in Table 8 and Figure 4. The table contains the AUC and OA mean values in both the test and train datasets. Moreover, it shows the mean CPU time needed to train each method. The figure analyzes the trade-off between generalization capability (test AUC) and computational cost (CPU train time). Finally, Table 9 shows the estimated specificity and sensitivity values.

Table 8: Results in the *sonar* semi-synthetic dataset. Test and train performances (in terms of AUC and OA) and the CPU time needed to train each method are provided. The results are the mean over the 10 runs. The best generalization (test) performance among the crowdsourcing methods that do not use test set annotations (central rows) is bolded.

Methods	Test set		Train set		CPU time (s)
	AUC	OA%	AUC	OA%	
<i>GP-GOLD</i>	0.9043	80.16	0.9901	94.32	98.09
<i>GP-MV</i>	0.7779	60.97	0.8822	71.51	34.05
<i>Rodrigues</i>	0.8574	71.77	0.9843	91.10	153.67
<i>VGPCR</i>	<b>0.8668</b>	<b>74.84</b>	0.9680	89.59	172.09
<i>Raykar</i>	0.6974	65.65	0.9115	88.08	52.65
<i>Yan</i>	0.6592	57.58	0.7698	75.41	228.72
<i>MV*</i>	0.8452	77.42	0.8449	77.60	0
<i>VGPCR*</i>	0.9890	94.19	0.9851	93.15	172.09

We observe again that *VGPCR* obtains the best generalization performance

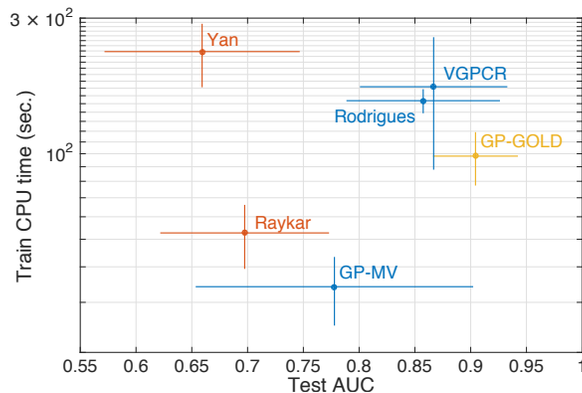


Figure 4: Trade-off between generalization capability (in terms of test AUC) and computational cost (in terms of CPU train time) in the *sonar* dataset. For each method, the mean plus/minus one standard deviation is shown. The color indicates the family of the algorithm, i.e. yellow for the non-crowdsourcing method *GP-GOLD*, blue for the GP-based crowdsourcing methods, and red for the LR-based ones. Notice also the logarithmic scale in the y-axis.

among the five crowdsourcing methods that do not use test set annotations (in both test AUC and test OA). Moreover, it clearly obtains the most accurate estimations of specificity and sensitivity: the maximum absolute difference in  
500 Table 9 is 0.0266 for *VGPCR*, whereas it is 0.1081 for *Raykar* and 0.3039 for *Rodrigues*. The training CPU time is similar to the one obtained by *Rodrigues* (the only crowdsourcing method that is competitive with it in test performance).

Let us analyze the results for the LR-based methods, which will again shed some light on the internal structure of the underlying classification dataset. As  
505 opposed to the *heart* dataset, here the test results for *Raykar* (and *Yan*) are distinctly worse than those for the GP-based methods (specially the more elaborated *VGPCR* and *Rodrigues*, see Figure 4, which clearly shows that blue points are to the right of red ones in the x-axis). This suggests that this *sonar* database is not as linearly-separable as the one before. Again, this can be confirmed by  
510 training a standard LR classifier with the true labels. Indeed, it obtains a mean test AUC value of 0.7546, very far from the more complex decision boundary of *GP-GOLD* (0.9043, see Table 8). Notice also that this LR classifier is an intu-

Table 9: Estimated values of sensitivity and specificity for the five annotators in the *sonar* semi-synthetic experiment. Only those methods that include these parameters in their formulation are shown. The values are the mean over the 10 realizations.

Annotator	Original		<i>Raykar</i>		<i>Rodrigues</i>		<i>VGPCR</i>	
	$\alpha$	$\beta$	$\alpha$	$\beta$	$\alpha$	$\beta$	$\alpha$	$\beta$
1	0.9	0.6	0.8889	0.7081	0.6768	0.4374	0.8734	0.6244
2	0.7	0.8	0.6786	0.8548	0.4890	0.6221	0.7027	0.8167
3	0.8	0.5	0.8215	0.5296	0.7003	0.3717	0.8051	0.4761
4	0.1	0.2	0.2054	0.2580	0.3922	0.4850	0.0961	0.2079
5	0.9	0.8	0.8359	0.8253	0.5961	0.5334	0.8815	0.7997

itive upper bound for the LR-based crowdsourcing methods (just as *GP-GOLD* is for the GP-based ones). This is in accordance with the test AUC values obtained by *Raykar* and *Yan* (0.6974 and 0.6592, respectively), which are below 0.7546. Regarding the comparison between them, *Yan* is again significantly outperformed by *Raykar*, which is also much faster. The justification is as before: *Yan*'s feature-dependent model is too complex for the simple generation process of the annotations, which follows the simpler model of *Raykar*. This makes the convergence more difficult for *Yan*, whereas *Raykar* logically gets pretty good estimations of  $\alpha$  and  $\beta$  (see Table 9).

The behavior of the GP-based methods is the expected one. Unlike in the *heart* dataset, where it suffered over-fitting, here *Rodrigues* (and also *VGPCR*) exhibit better predictive performance than their intuitive lower bound *GP-MV*. They are also upper bounded by their natural limit *GP-GOLD*. Interestingly, we see that the difference here between *GP-GOLD* and *GP-MV* (in test AUC and OA) is significantly larger than in *heart*. This is connected with the aforementioned non-linearity of the dataset: a close-to-linear boundary can be well identified with low-quality labels, but a complex one needs more accurate data.

The two GP-based methods *VGPCR* and *Rodrigues* present a very similar trade-off between predictive performance and computational cost (Figure 4). However, the estimations of  $\alpha$  and  $\beta$  are much poorer for *Rodrigues*. In principle

this is certainly surprising, because the formulas that define  $\alpha$  and  $\beta$  in this work (recall eqs. (11)-(12)) are the same as in *Rodrigues* (see eqs. (8)-(10) in [12])<sup>4</sup>.  
 535 There are two explanations for this: 1) the treatment of the latent variable  $\mathbf{z}$  (which appears in the formulas for  $\alpha$  and  $\beta$ ), and 2) the modeling of  $\alpha$  and  $\beta$  themselves. The first one is pretty subtle but very relevant, and refers to the fact that  $\mathbf{z}$  is integrated out from the beginning in the model of [12] whereas it is included in our model as a latent variable. That allows us to compute  
 540 sounder estimates for  $\mathbf{z}$ , which is the basis of the  $\alpha$  and  $\beta$  update formulas. The second one is clearer, as our posterior distributions over  $\alpha$  and  $\beta$  account for the uncertainty in the model (whereas the point estimates in [12] do not).

The conclusions for the methods that make use of the test annotations is the same as in the *heart* dataset: *VGPCR\** obtains extraordinarily good results  
 545 thanks to the accurate estimation of  $\alpha$  and  $\beta$ , whereas *MV\** is not competitive with it because it does not model the noise in the annotations.

### 5.3. Fully real data

In this section we compare the performance of *VGPCR* and its competitors on two real crowdsourcing datasets. True labels for the training instances are  
 550 provided by the datasets contributors. This allows us to compare also with *GP-GOLD*. However, no test annotations  $\mathbf{y}_*$  are provided, so *VGPCR\** and *MV\** are not included in this section. The obtained results support that the novel *VGPCR* is also the most competitive approach in these practical applications.

#### 5.3.1. Sentence Polarity dataset

555 The *Sentence Polarity* dataset first was presented by Pang and Lee [34]. It consists of 10427 sentences extracted from movie reviews in “Rotten Tomatoes” website <http://www.rottentomatoes.com/>. The goal is to decide whether a sentence corresponds to a “positive” or “negative” review. In Table 10 we

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<sup>4</sup>More precisely, recall that we model  $\alpha$  and  $\beta$  as stochastic variables whereas they are treated as parameters in [12]. Thus, it is the *mean* of the beta distributions in eqs.(11)-(12) what equals the formulas in [12].

Table 10: Examples of positive and negative samples in *Sentence Polarity* dataset.

Sentence	True Label
“An original gem about an obsession with time.” “A taut, intelligent psychological drama.” “Clever, brutal and strangely soulful movie.”	“positive”
“This is amusing for about three minutes.” “The film can depress you about life itself.” “The pool drowned me in boredom.”	“negative”

show six sentences in the dataset. Preprocessing and feature extraction were  
 560 carried out by Rodrigues *et al.* [35], which resulted in feature vectors with 1200  
 components. The dataset is divided into train and test sets, with 4999 and 5428  
 samples, respectively. To obtain crowdsourcing labels, the train set was made  
 available in Amazon Mechanical Turk. A total amount of 27746 labels were  
 obtained from 203 different annotators.

565 Results are shown in Table 11 and Figure 5. The table contains the AUC and  
 OA for both test and train datasets. Moreover, it shows the CPU time needed  
 to train each method. The figure analyzes the trade-off between generalization  
 capability (test AUC) and computational cost (CPU train time). Finally, Figure  
 6 shows the estimated specificity and sensitivity values.

570 Again, *VGPCR* is the best crowdsourcing method in terms of predictive  
 performance (0.8 of AUC and 72.53% of OA). These values place the proposed  
 method really close to its natural upper bound *GP-GOLD* (0.8037 of AUC and  
 73.07% of OA). As expected, it is also lower bounded by *GP-MV*.

It is important here to analyze the behavior of *Rodrigues*. Although it is  
 575 below its intuitive lower bound *GP-MV* (it is clearly suffering from over-fitting,  
 see its high training performance), its generalization capability is not very far  
 from *VGPCR*, and it seems that it might be the method of choice in certain  
 applications because it is around four (resp. three) times faster than *VGPCR*  
 (resp. *GP-MV*). This low computational cost (the lowest in this dataset) seems  
 580 certainly surprising, since the EP inference is quite expensive (as both semi-

Table 11: Results in the *Sentence Polarity* fully real dataset. Test and train performances (in terms of AUC and OA) and the CPU time needed to train each method are provided. The results are the mean over the 10 runs. The best generalization (test) performance among the crowdsourcing methods is bolded.

Methods	Test set		Train set		CPU time (s)
	AUC	OA%	AUC	OA%	
<i>GP-GOLD</i>	0.8037	73.07	0.9130	83.76	$3.3089140 \times 10^4$
<i>GP-MV</i>	0.7932	72.03	0.8706	79.22	$2.9595670 \times 10^4$
<i>Rodrigues</i>	0.7815	72.07	0.9415	89.44	$1.0685530 \times 10^4$
<i>VGPCR</i>	<b>0.8000</b>	<b>72.53</b>	0.8861	81.32	$3.9638080 \times 10^4$
<i>Raykar</i>	0.7141	68.22	0.9100	90.68	$1.8156210 \times 10^4$
<i>Yan</i>	0.7530	69.45	0.8974	84.28	$1.4089233 \times 10^5$

synthetic experiments have shown). The key is that, in this application, the code provided by the authors fixes the kernel hyperparameters from the beginning<sup>5</sup>, and they are not estimated during training (which is the most time-consuming step). However, the other two GP-based methods do estimate them. Therefore, the CPU training costs should not be compared. If we fix the kernel hyperparameters of *VGPCR* to its previously estimated values, then its CPU training time falls down to  $2.1312 \times 10^3$  seconds (around 5 times less than *Rodrigues*), whereas its predictive performance remains unchanged.

The problem is that, because of the EP inference procedure, estimating the kernel hyperparameters with *Rodrigues* in this dataset (4999 training instances) is computationally prohibitive. Indeed, Figure 7 shows the experimental CPU time needed to train *Rodrigues* (including the hyperparameters estimation) with increasingly larger subsets of the original set. The rapid growth makes training with  $n = 4999$  instances infeasible. Interestingly, the theoretical complexity of each EP iteration is  $\mathcal{O}(n^3)$ , the same as the variational inference used here. However, EP usually requires many more iterations for convergence, and that

<sup>5</sup>Specifically, the length-scale  $l$  is fixed to 1.5 and the variance  $\gamma$  to 1.3 (recall Section 2).

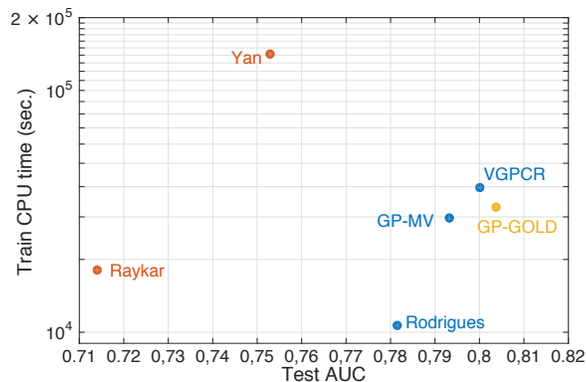
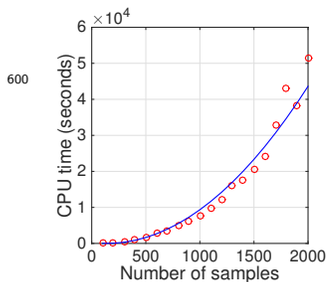


Figure 5: Trade-off between generalization capability (test AUC) and computational cost (CPU train time) in the *Sentence Polarity* dataset. The color denotes the family of the algorithm: yellow for the non-crowdsourcing method *GP-GOLD*, blue for the GP-based crowdsourcing methods, and red for the LR-based ones. Notice the logarithmic scale in the y-axis.

makes it computationally heavier in practice.



605 Figure 7: Mean CPU time in three runs of *Rodrigues*. The x-axis shows the number of training samples.

Regarding the estimation of specificity and sensitivity, Figure 6 shows very similar estimations for *VG-PCR* and *Raykar*, whereas *Rodrigues* deviates from this common tendency. This is in accordance with all the previous experiments, where *Rodrigues* estimations were less reliable. The reasons behind this were analyzed in Section 5.2.2.

Figure 5 shows a clear separation between GP- and LR-based methods in the x-axis (i.e., the generalization capability). As explained in Section 5.2, this may reveal a non-linear underlying structure in the dataset.

610 Finally, as opposed to the semi-synthetic datasets, notice that *Yan* significantly outperforms *Raykar* here. This is in accordance with the fact that the annotations generation process does not necessarily imitate *Raykar*'s one in this real dataset, and the feature-dependent model of *Yan* seems to adapt well.

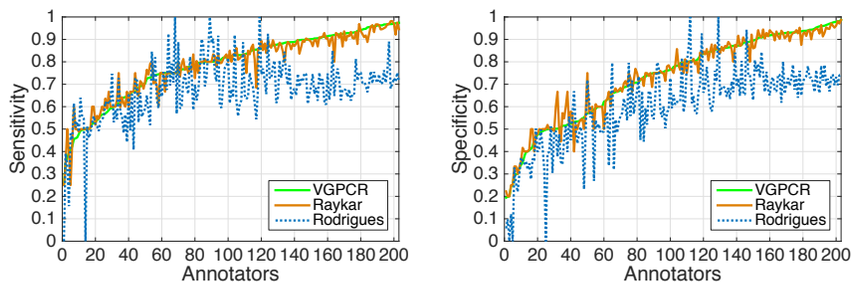


Figure 6: Sensitivity  $\alpha$  (left) and specificity  $\beta$  (right) estimations for the 203 annotators in the *Sentence Polarity* dataset. Only those methods that include these parameters in their formulation are shown. For a clearer display, in each figure the annotators are arranged in ascending order of the *VGPCR* estimated value.

However, this is at the expense of a really heavy training step, being the only  
 615 method (together with Rodrigues) beyond  $10^5$  seconds of CPU train time.

### 5.3.2. Music Genre dataset

In this experiment we use the Music Genre dataset presented in [36], which consists of 1000 fragments (30 secs. length) of songs. The goal is to distinguish between 10 music genres: *classical*, *country*, *disco*, *hiphop*, *jazz*, *rock*, *blues*,  
 620 *reggae*, *pop*, and *metal*. We use an *one-vs-all* strategy to address this multi-class classification problem, and the results are averaged over the 10 experiments.

For preprocessing and feature extraction, the authors in [35] used Marsyas music information tool (<http://marsyas.info/>) to extract 124 features from the original dataset. These features include relevant technical metrics such as  
 625 means and variances of timbral features, time-domain zero-crossings, spectral centroid, rolloff, flux, and Mel-Frequency Cepstral Coefficients (MFCC).

The dataset contains 100 samples from each genre, which were randomly divided in 70 samples for training and 30 for testing. Crowdsourcing labels were obtained with Amazon Mechanical Turk. Each annotator listened to a  
 630 subset of fragments and labeled them as one of the ten genres listed above. A total amount of 2945 labels were provided by 44 different annotators.

The results are shown in Table 12 and Figure 8. The table contains the

Table 12: Results in the *Music Genre* fully real dataset. Test and train performances (in terms of AUC and OA) and the CPU time needed to train each method are provided. The results are the mean over the 10 runs. The best generalization (test) performance among the crowdsourcing methods is bolded.

Methods	Test set		Train set		CPU Time (s)
	AUC	OA%	AUC	OA%	
<i>GP-GOLD</i>	0.9426	94.60	0.9713	95.69	$3.283342 \times 10^3$
<i>GP-MV</i>	0.8865	91.50	0.8809	91.89	$2.170080 \times 10^3$
<i>Rodrigues</i>	0.8795	85.43	0.9429	92.16	$6.520268 \times 10^3$
<i>VGPCR</i>	<b>0.9152</b>	<b>92.70</b>	0.9259	93.70	$1.712601 \times 10^3$
<i>Raykar</i>	0.8806	90.40	0.9414	95.84	$7.201810 \times 10^2$
<i>Yan</i>	0.8614	91.90	0.8913	93.96	$1.088944 \times 10^3$

AUC and OA for both test and train datasets. Moreover, it shows the CPU time needed to train each method. The figure analyzes the trade-off between generalization capability (test AUC) and computational cost (CPU train time).  
 635 Finally, Figure 9 shows the estimated specificity and sensitivity values.

Once more, the novel *VGPCR* exhibits the best generalization capability, keeping a considerable distance with the next one (*GP-MV*). Moreover, *VGPCR* is also the fastest among the GP-based methods. This implies an unbeatable trade-off in Figure 8. Furthermore, as theoretically expected, its performance  
 640 lies between that of *GP-GOLD* and *GP-MV*.

As opposed to the previous experiment, *Rodrigues* is now the most computationally expensive method (around three times more than the next one, *GP-MV*). This difference is due to the fact that the kernel hyperparameters are estimated during the training step. Test performance for *Rodrigues* is clearly  
 645 below *VGPCR*, being only competitive with *GP-MV*. This is due to over-fitting (see its high training performance in comparison with the test one), and the very poor estimation of sensitivity/specificity (see Figure 9). In turn, as explained in Section 5.2, these follow from a less subtle modeling (marginalization of  $\mathbf{z}$ ,  
 650 point estimates for  $\boldsymbol{\alpha}$  and  $\boldsymbol{\beta}$ ) and the use of a different inference approach.

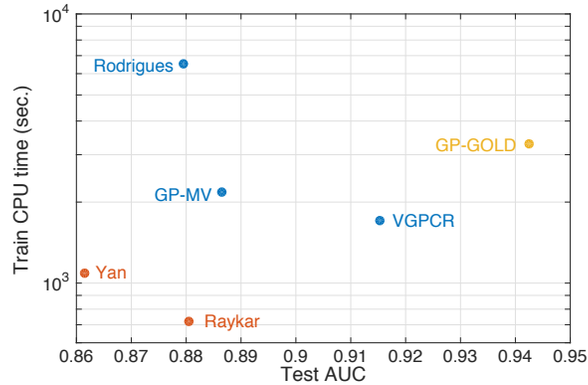


Figure 8: Trade-off between generalization capability (test AUC) and computational cost (CPU train time) in the *Music Genre* dataset. The color indicates the family of the algorithm: yellow for the non-crowdsourcing method *GP-GOLD*, blue for the GP-based crowdsourcing methods, and red for the LR-based ones. Notice the logarithmic scale in the y-axis.

Once more, Figure 9 shows similar estimates for *VGPCR* and *Raykar*, whereas *Rodrigues* exhibits a quite bizarre behavior with almost constant estimates. This definitely confirms its difficulties for calculating  $\alpha$  and  $\beta$ . In this particular case, the problem may come from the unbalanced setting (recall the *one-vs-all* strategy, which implies a 90%-10% balance between negative and positive classes). In fact, *Rodrigues* is the only method with test OA below 90%, which would be the OA for a naive classifier that assigns every instance to the majority class.

In the comparison between GP- and LR-based methods, the x-axis of Figure 8 does not show a clear separation in predictive performance. This suggests that linear boundaries may be representative for the classes of this set. Indeed, notice that both families are much better separated in the y-axis. Interestingly, this is precisely connected with the aforementioned underlying linear structure, which allows for a fast convergence of the LR-based methods. The same behavior could be appreciated in the close-to-linear *heart* set, recall Figure 3.

Regarding the LR-based methods, *Raykar* and *Yan* obtain similar results (the former performs better with respect to AUC and the latter with respect to OA). However, the complex modeling of *Yan* makes it computationally heavier,

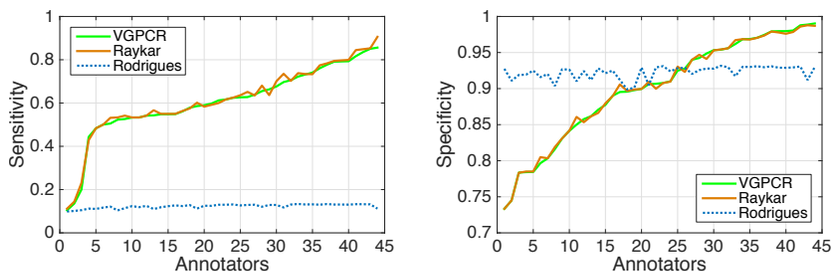


Figure 9: Sensitivity  $\alpha$  (left) and specificity  $\beta$  (right) estimations for the 44 annotators in the *Music Genre* dataset. Only those methods that include these parameters in their formulation are shown. For a clearer display, in each figure the annotators are arranged in ascending order of the *VGPCR* estimated value.

and thus less competitive in practice.

## 6. Conclusions

670 We have introduced a new crowdsourcing classification methodology. As  
previous approaches, it is based on a Gaussian Process classifier, which allows  
for the description of complex data. However, a novel Variational Bayes (VB)  
inference procedure is proposed here (instead of Expectation Propagation, EP).  
The modeling of the annotators is also refined with respect to previous GP-  
675 based methods: the level of expertise is treated as a stochastic variable, and  
the underlying true training labels  $\mathbf{z}$  are not marginalized out from the model.  
Moreover, the proposed method allows for integrating in the prediction (possibly  
non-expert) annotations that may have been provided for test instances.

The experimental results have shown that the novel VB-based approach is  
680 really competitive and robust across very different types of datasets, ranking  
always first among its competitors in terms of predictive performance. On the  
contrary, the EP-based method has suffered over-fitting in three out of the five  
datasets used. The computational cost of the proposed method is competitive  
with the rest of the crowdsourcing classifiers (and considerably lower than the  
685 EP-based one). Our refined model for the annotators is also reflected in the ex-  
periments. Indeed, our sensitivity-specificity estimations are significantly more

accurate than those by the EP-based approach. If there are test annotations available (which is not always possible, see the fully real datasets used here), we have seen that the proposed method largely benefits from its probabilistic integration within the model. It would be interesting to study to what extent this generalizes to fully real datasets, where the annotations generation process does not necessarily follow the one proposed in the model. Other lines of future work are i) development of alternative and more accurate feature-dependent crowdsourcing models, ii) a probabilistic multi-class generalization of the proposed model, and iii) extension of GP-based crowdsourcing methods to large-scale datasets.

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