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Cost and Quality Assurance in Crowdsourcing Workflows*

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ABSTRACT

Crowdsourcing platforms provide tools to replicate and distribute micro tasks (simple, independent work units) to the crowd and assemble results. However, real-life problems are often complex: they require to collect, organize, process or transform data, guarantee the quality of results, and meet budget constraints. Further, specifications and realization of intricate jobs on crowdsourcing platforms are still in their infancy.

In this work, we combine workflows with crowdsourcing to enable efficient execution of complex tasks. Workflows provide ways to organize a complex task in phases and guide the overall realization. The challenge is to interface workflows and crowdsourcing systems efficiently to achieve good accuracy of results at a reasonable cost. Standard "static" allocation of work in crowdsourcing affects a fixed number of workers per micro-task to realize and aggregates the results. We propose synchronous and asynchronous dynamic worker allocation techniques on top of workflows, where decisions to replicate tasks, worker allocation and progress of the workflow execution depend on inferred tasks difficulty, worker expertise, confidence in answers and remaining budget. We evaluate the performance of this framework on a benchmark and show that the proposed approaches outperform static allocation in terms of cost and accuracy.

KEYWORDS

Crowdsourcing; Data-centric workflows

1 INTRODUCTION

Despite recent advances in artificial intelligence and machine learning, many tasks still require human contributions. With the growing availability of internet, it is now possible to hire workers all around the world on crowdsourcing marketplaces. Many crowdsourcing platforms have emerged in the last decade: Amazon Mechanical Turk¹, Figure Eight², Wirk³, etc. They hire workers from a crowd to solve problems [25]. A platform allows employers to post tasks, that are then realized by workers in exchange for some incentives [6]. Common tasks include image annotation, surveys, classification, recommendation, sentiment analysis, etc. [12]. The existing platforms support simple, repetitive and independent *micro-tasks* which require few minutes to an hour to complete.

However, many real-world problems are not simple micro tasks, but rather complex orchestrations of dependent tasks, which aims are to process input data and workers answers. The existing crowdsourcing platforms provide interfaces to execute micro tasks and access crowd, but lack ways to specify and execute complex tasks. The next stage of crowdsourcing is to design systems to realize more complex and involved tasks over existing

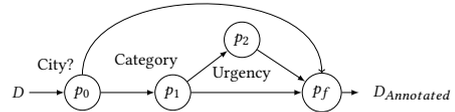


Figure 1: A workflow in a smart city

crowdsourcing platforms. A natural solution is to define complex tasks as workflows, i.e., orchestrations of jobs that exchange data to achieve a final objective [30]. The output of individual jobs (documents, annotations, data) is passed to the next tasks according to the workflow rules.

Complex workflows appear in many contexts. Consider for instance online request services frequently proposed by smart cities (Figure 1). These systems receive a huge number of requests that must be filtered, classified and then sent to the appropriate bureau of the city. This requires a lot of work power, but it follows a workflow made of several simple stages. A first step p_0 sorts a load of demands D according to the address of the client (demands from residents of the city will not be processed like external demands). The demands from city inhabitants are validated and sent to the next phase p_1 . The goal of p_1 is to decide in which category (Health, roads, ...) the request falls. After classification, phase p_2 decides whether the demand is *Urgent* or not if some demands from that category call for rapid response. For categories that are not urgent, demands are forwarded directly to p_f . The final phase p_f gathers all annotated demands in an annotated dataset $D_{Annotated}$.

Workflows alone are not sufficient to handle complex tasks with crowdsourcing. Many data-centric applications come with budget and quality constraints: As human workers are prone to errors, one has to hire several workers to aggregate a final answer with sufficient confidence. An unlimited budget would allow hiring large pools of human workers to assemble reliable answers for each micro-task, but in general, a client for a complex task imposes a limited budget B_0 that must not be exceeded. A limited budget forces to replicate micro-tasks in an optimal way to achieve the best possible quality. The objective is then to obtain a *reliable* result, forged through a complex orchestration, and at a *reasonable cost*.

In this paper, we propose a solution for the efficient realization of complex tasks. We define a workflow model, which orchestrates tasks and work distribution according to a dynamic policy that considers confidence in aggregated data and the cost to increase this confidence. A workflow can be seen as an orchestration of phases, where the goal of each phase is to tag records from the dataset input to the phase. A complex task terminates when the last of its phases has completed its tagging. The output of one phase acts as an input to the next ones in the workflow. For simplicity, we consider simple Boolean tagging tasks that associate a tag in $\{0, 1\}$ to every record in a dataset. We also assume that workers are uniformly paid. As humans are prone to errors, tagging must be performed by several workers, and the answers assembled with aggregation technique. For each record, one of the possible answer (called the *ground truth*) is correct, and an aggregated answer is considered as reliable if its probability to

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¹www.mturk.com, ²www.appen.com, ³www.wirk.com

be the ground truth is high. Hiring more workers to tag records increases the reliability of the aggregated answer. The overall challenge is hence to realize a workflow within budget B_0 , while guaranteeing that the final dataset forged during the last phase of the workflow has a high probability to be the ground truth.

This general orchestration schema leaves several design choices open. First, the aggregation technique used influences the quality of the final results. Furthermore, the mechanisms used to decide if more workers must be hired to improve quality have an impact on costs and on the overall accuracy of answers. The simplest way to replicate micro tasks is static execution, i.e. affect an identical fixed number of workers to each micro-task in the orchestration without exceeding budget B_0 . On the other hand, one can allocate workers to tasks *dynamically*. One can wait in each phase to achieve a sufficient reliability of answers for each record of the input before forwarding data. This is called a *synchronous* execution of a workflow. Last, one can eagerly forward records with reliable tags to the next phases without waiting for the total completion of a phase. This is called *asynchronous* execution.

In this work, we study execution strategies for complex workflows with different design choices. We consider several types of workflow, of aggregation (namely Majority Voting (MV) and Expectation Maximization (EM) based techniques [13]) several distributions of data, difficulty of tasks and workers expertise, and study the cost and accuracy of workflows execution under static, synchronous and asynchronous assignment of workers to tasks. Unsurprisingly, dynamic distribution of work allows to save costs in all cases. A more surprising result is that synchronous realization of complex tasks is in general more efficient than asynchronous realization.

Related Work: Several works have considered complex workflows, interactions with crowdsourcing systems, or advanced aggregation techniques to improve data quality. We do not claim exhaustiveness but list below some papers which are the closest to our complex crowdsourcing solution. Coordination of tasks has been considered in many languages such as BPMN [24], ORC [16, 21], BPEL [2, 23], or workflow nets[31], a variant of Petri nets dedicated to business processes. They allow parallel or sequential execution of tasks, fork and join operations to create or merge a finite number of parallel threads. Tasks are represented by transitions. Workflow nets mainly deal with the control part of business processes, and data is not central for this model: Tasks are not replicated, and token do not carry data. The more advanced versions of the model [10] handle data through global variables. Similarly, BMPN, ORC and BPEL are centered on the notion of transaction, and are not tailored for datasets manipulations, replication of tasks, nor aggregation of results. To overcome the limitations of orchestration models, several data-centric models have emerged. Guarded Active XML [1] is a specification model where references to external services are introduced in structured data. Services can modify data when their guard is satisfied, and replace a part of the data by some computed value that may contain new references to service calls. Business artifacts were originally developed by IBM [22]. Artifacts and their many variants are data-centric, i.e. they focus on possible modifications of datasets within a system. [14] considers Data-Centric Dynamic Systems (DCDS), i.e. relational databases equipped with guarded actions that can modify their contents, and call external services. Most of these data-centric models are Turing powerful, and hence in their full generality can be seen as programs coordinating data transformations (or calls to a crowdsourcing platform). Their strength lies in their capacity to describe formally how data is

transformed, and for some models in the existence of subsets of the language that can be formally verified [1, 5]. However, they are tailored to manipulate data within a single transaction, and do not consider replication of tasks nor aggregation of results, which are essential in crowdsourcing.

Some works propose empirical solutions for complex data acquisition, mainly at the level of micro-tasks [12, 19]. Crowdforge uses Map-Reduce techniques along with a graphical interface to solve complex tasks [17]. Turkit [20] is a crash and rerun programming model. It builds on an imperative language, that allows for repeated calls to services provided by a crowdsourcing platform. Turkomatic [18] is a tool that recruits crowd workers to help clients planning and solving complex jobs. It implements a Price, Divide and Solve (PDS) loop, that asks crowd workers to divide a task into orchestrations of subtasks, and repeats this operation up to the level of micro-tasks. A PDS scheme is also used by [33] in a model based on hierarchical state machines that orchestrates sub-tasks.

In this work, we assemble answers returned by workers using aggregation techniques. Basic aggregation uses majority voting (MV), i.e. takes as final result for a tagging task the most returned answer. Several approaches have improved MV by using Expectation Maximization (EM) or by considering workers competences, expressed in terms of accuracy (ratio of correct answers) or in terms of recall and specificity (that considers correct classification for each possible type of answer). It is usually admitted [34] that recall and specificity give a finer picture of worker's competence than accuracy. We only highlight works that focus on EM or MV to aggregate data, and refer interested readers to [34] for a more complete survey of the domain. Zencrowd [8] considers workers competences in terms of accuracy and aggregates answers using EM. Workers accuracy and ground truth are the hidden variables that must be discovered in order to minimize the deviations between workers answers and aggregated conclusion. D&S [7] uses EM to synthesize answers that minimize error rates from a set of patient records. It considers recall and specificity, but not the difficulty of tasks. [15] proposes an algorithm to assign tasks to workers, synthesize answers, and reduce the cost of crowdsourcing. It assumes that all tasks have the same difficulty, and that workers reliability is a static probability to return the correct value that applies to all types of tasks. EM is used by [27] to discover recall and specificity of workers and propose maximum-likelihood estimator that jointly learns a classifier, discovers the best experts, and estimates ground truth. Most of the works cited above consider expertise of workers but do not address tasks difficulty. Approaches such as GLAD [32] or [4] also estimate tasks difficulty to improve quality of answers aggregation on a single dataset.

Generally the database and machine learning communities focus on data aggregation techniques and leave budget optimization apart. CrowdBudget [29] is an approach that divides a budget B among K existing tasks to replicate them and then aggregate answers with MV. Raykar et.al [26] use Markov Decision Processes to decide whether answers to a database tagging scheme are sufficiently accurate or new worker should be hired. Crowdinc [28] is an EM-based aggregation technique that considers task difficulty, recall and specificity of workers. It computes accuracy of an aggregation, and launches new tasks dynamically to achieve a given threshold. The model proposed in this paper is a workflow that orchestrates tasks, replicates them, distributes them and aggregates the returned results before passing the forged dataset to the next tasks. It is a variant of the complex workflow

model proposed in [3], and it uses the aggregation technique of Crowdinc [28] to forge reliable answers.

The rest of the paper is organized as follows. Section 2 describes the complex workflow model, and Section 3 the aggregation technique used to forge answers from crowdworkers returns. In Section 4, we propose algorithms to realize complex workflows while maintaining a trade-off between cost and quality. We report the experimental evaluation and result in Section 5 before conclusion. Due to lack of space, some technical details are provided in a separate appendix at the end of the paper.

2 COMPLEX WORKFLOW WITH AGGREGATION

In this section, we formalize the complex workflow model. The model is inspired by data centric workflows [3], but adds tasks replication to this model, and further considers aggregation and budget management in its semantics. The context of the workflow is the following: A client wants to realize a complex task that needs the knowledge and skills of human workers. Complex tasks are divided into several dependent phases. Each phase processes records from an input dataset or merges different inputs to a single one, and forwards the result to its successor. Datasets are collections of records, i.e. relations of the form $r(v_1, \dots, v_k)$ where v_1, \dots, v_k are values for the fields of the record. One can use First Order statements to address properties of a record (e.g. write $v_i == true$), or of a set of records in a dataset (e.g. $\exists r(v_1, \dots, v_k) \in D, v_k == true$). We will denote by FO^R the FO formulas for records, and by FO^D the FO formulas for datasets. For simplicity, we assume that processing a record is a micro-task that simply consists in adding a new Boolean field (called a *tag*) to this record. Hence a micro-task can be seen as an operation that transforms a record $r(v_1, \dots, v_k)$ into a new record $r'(v_1, \dots, v_k, v_{k+1})$ where v_{k+1} is a Boolean value. This setting can be easily adapted to let v_{k+1} take values from a discrete domain.

As humans are prone to errors, micro-tasks are replicated, and allocated to several workers. Their answers are then aggregated before proceeding to the next tasks. Hence, an aggregation mechanism is required to combine the answers and forward the results to the next phases. When a phase has several successors, the contents of records is used to decide to which successor(s) it should be forwarded. This allows to split datasets according to the value of a particular data field, process differently record depending on their contents, create concurrent threads, etc. We define a workflow as follows.

Definition 2.1 (Complex Workflow). A complex workflow is a tuple $W = (\mathcal{P}, \longrightarrow, G, \otimes, p_0, p_f)$ where \mathcal{P} is a finite set of phases, p_0 is a particular phase without predecessor, p_f a phase without successor, $\longrightarrow \subseteq \mathcal{P} \times FO^R \times \mathcal{P}$ is a flow relation and $G = \mathcal{P} \rightarrow FO^D$ associates a guard to every phase, and for every $p_x \in \mathcal{P}$, $\otimes(p_x)$ is an operator used to merge input datasets.

Intuitively, a phase associates a tagging task to each record in a dataset, replicates and distributes it to several workers. The answers returned by all the workers are then aggregated to get a final trusted answer. We assume that workers answers are independent. For a triple $(p_x, g_{x,y}, p_y)$ in \longrightarrow , we will say that p_x is a predecessor of p_y . We denote by $Succ(p_x) = \{p_y \mid p_x \longrightarrow^* p_y\}$ the set of phases that must occur after p_x , i.e. and by $Pred(p_x) = \{p_y \mid p_y \longrightarrow^* p_x\}$ the set of phases that must occur before p_x . The meaning of guard $g_{x,y}$ is that every record produced by phase p_x that satisfies guard $g_{x,y}$ is forwarded to p_y . We will see in the

rest of this section that "producing a record" is not done in a single shot, and requires to duplicate a tagging micro-task, aggregate answers, and decide if the confidence in the aggregated answer is sufficient. When a phase p_x has several successors p_y^1, \dots, p_y^k and the guards $g_{x,y^1}, \dots, g_{x,y^k}$ are exclusive, each record processed by p_x is sent to at most one successor. We will say that p_x is an *exclusive fork* phase. On the contrary, when guards are not exclusive, a copy of each record processed in p_x can be sent to each successor (hence increasing the size of data processed in the workflow), and p_x is called a *non-exclusive fork* phase. For a given phase $p_x \in \mathcal{P}$, we denote by $G_x \in FO^D$ the guard attached to phase p_x . G_x addresses properties of the datasets input to p_x by its predecessors. This allows in particular to require that all records in preceding phases have been processed (we will then say that phase p_x is *synchronous*), that at least one record exists in some predecessor (the task is then fully *asynchronous*), or any FO expressible property on datasets produced by predecessors of p_x . Similarly, we denote by $\otimes_x = \otimes(p_x)$ the operator used to merge all inputs entering phase p_x . This operator can be either a simple union of datasets, or a more complex join operation. If \otimes_x is a join operation, we impose that p_x is synchronous. This is reasonable, as one cannot start processing data produced by a join operation when the final set of records is not known. When \otimes_x is a simple union of datasets, as tasks are individual tagging of records, any record processed on a predecessor of p_x can be processed individually without waiting for other results to be available. This allows asynchronous executions in which two phases p_x, p_y can be concurrently active (i.e. have started processing records), even if p_x precedes p_y . On the contrary, if the execution of a phase p_x is synchronous, and p_y is a successor of p_x all records output to p_y must be tagged before starting phase p_y .

The semantics of a complex workflow is defined in terms of moves from a configuration to the next one, organized in *rounds*. Configurations memorize the data received by phases, a remaining budget, the answers of workers, and quality measures on answers and workers competences.

Definition 2.2. A configuration of a complex workflow is a tuple $C = (\mathcal{D}_{in}, W_{in}, W_{out}, conf, exp, B)$ where

- $\mathcal{D}_{in} : \mathcal{P} \rightarrow Dsets$ Associates a dataset to every phase $p_x \in \mathcal{P}$ (this dataset can be empty).
- $W_{in} : \mathcal{P} \times \mathbb{N} \times \mathbb{N} \rightarrow 2^W$ associates a set of workers to each record in $\mathcal{D}_{in}(p_x)$.
- $W_{out} : \mathcal{P} \times \mathbb{N} \times W \rightarrow \{0, 1\} \cup \emptyset$ associates a tag or the empty set to a worker, a phase and a record number that was formerly affected to the worker. We will write $l_{i,j}^x$ to denote the answer returned by worker w_i when tagging record r_j during phase p_x .
- $conf : \mathcal{P} \times \mathbb{N} \rightarrow [0, 1]$ is a map that associates to each record in $\mathcal{D}_{in}(p_x)$ a confidence score in $[0, 1]$ computed from W_{out}
- $exp : W \times \mathcal{P} \rightarrow [0, 1]$ is a confidence score that associates to each worker a score between 0 and 1.
- $B_r \in \mathbb{N}$ is the remaining budget.

$W_{out}(p_x, k, w) = \emptyset$ indicates that a worker w in a phase p_x has not yet processed record r_k . We say that phase p_x is *completed* for a record r_k from $\mathcal{D}_{in}(p_x)$ if there is no worker w such that $W_{out}(p_x, w, r_k) = \emptyset$. As soon as phase p_x is completed for r_k , we can derive an *aggregated answer* $r'_k(v_1, \dots, v_n, y_k^x)$ for each record $r_k(v_1, \dots, v_n)$ from the set of all answers returned by the workers in $W_{in}(p_x, k)$. Similarly, we can compute a confidence score

$conf(p_x, k)$ on the value y_k^x and the expertise of each worker (we will see how these values are evaluated in section 4). We say that a record r_i in a phase p_x is *inactive* if no more workers are assigned to it. It is *active* otherwise. Given a threshold value Th , will say that p_x is *finished* for a record r_k from $\mathcal{D}_{in}(p_x)$ if p_x is completed and $conf(p_x, k) > Th$. Record $r'_k(v_1, \dots, v_n, y_k^x)$ will then be part of the input of phase p_y if $(p_x, g_{x,y}, p_y) \in \rightarrow$ and $r'_k(v_1, \dots, v_n, y_k^x)$ satisfies guard $g_{x,y}$.

We can now detail how rounds change the configuration of a workflow. The key idea is that each round aggregates available answers, and then decides whether the confidence in aggregated results is sufficient. If confidence in a record is high enough, this record is forwarded to the successor phases, if not new workers are hired for the next round, which decreases the remaining budget. The threshold for the confidence decreases accordingly. Then new workers are hired for freshly forwarded data, leaving the system ready for the next round. From a configuration $C = (\mathcal{D}_{in}, W_{in}, W_{out}, conf, exp, B)$, a round produces a new configuration $C' = (\mathcal{D}_{in}, W_{in}, W_{out}, conf, exp, B)$ as follows:

- **Answers:** Workers that were hired in preceding round produce new data. For every phase p_x every record $r_n \in \mathcal{D}_{in}(p_x)$ and every worker w_i such that $w_i \in W_{in}(p_x, n)$ and $W_{out}(p_x, w_i, n) = \emptyset$, we produce a new output $l_{i,n}^x \in \{0, 1\}$ and set $W_{out}(p_x, w_i, n) = l_{i,n}^x$.
- **Aggregation:** The system evaluates aggregated answers in every active phase p_x . For every record r_k in $\mathcal{D}_{in}(p_x)$, we compute an aggregated answer y_n^x from the set of answers $A_n = \{l_{i,n}^x \mid w_i \in W_{in}(p_x, n)\}$. We also compute a new confidence score $conf'(p_x, n)$ for the aggregated answer (this confidence depends on the aggregation technique), and evaluate tasks difficulty and workers expertise (with the algorithm shown in Section 3).
- **Data forwarding :** We distinguish asynchronous and synchronous phases. Let p_y be an asynchronous phase (\otimes_y can only be a union of records). Then p_y accept every new record $r'(v_1, \dots, v_k, y_n^x)$ that was not yet among its inputs from a predecessor p_x provided r' satisfies guard $g_{x,y}$, and the confidence in the aggregated answer y_n^x is high enough. Formally, $\mathcal{D}'_{in}(p_y) = \mathcal{D}_{in}(p_y) \cup \{r'(v_1, \dots, v_k, y_n^x)\}$ if $(p_x, g_{x,y}, p_y) \in \rightarrow$, $conf'(p_x, n) \geq Th$ and $r'(v_1, \dots, v_k, y_n^x) \models g_{x,y}$. Let p_y be a phase such that \otimes_y is synchronous. We will say that a phase is closed if all its predecessors are closed, and for every $n, r_n \in \mathcal{D}_{in}(p_x)$, $conf(p_x, n) \geq Th$. If there exists a predecessor p_x of p_y that is not closed, then $\mathcal{D}'_{in}(p_y) = \emptyset$. Otherwise we can compute an input for phase p_y as a join over datasets computed by all preceding phases. Formally,

$$\mathcal{D}'_{in}(p_y) = \bigotimes_{\substack{y \\ p_x \rightarrow p_y}} D_x$$

where p_x ranges over the set of predecessors of p_y , and

$$D_x = \{ r(v_1, \dots, v_k, y_n^x) \mid r(v_1, \dots, v_k) \in \mathcal{D}_{in}(p_x) \\ \wedge r(v_1, \dots, v_k, y_n^x) \models g_{x,y} \}$$

Hence, for synchronous a phase p_y , the input dataset is a join operation computed over datasets filtered by guards, and realized only once predecessor tasks have produced all their results. Both in synchronous and asynchronous settings, phase p_y becomes active if $\mathcal{D}'_{in}(p_y) \models G(p_y)$, and we set $conf'(p_y, n) = 0$ for every new record in $\mathcal{D}'_{in}(p_y)$

- **Worker allocation :** For every p_x that is active and every record $r_n = r(v_1, \dots, v_k) \in \mathcal{D}'_{in}(p_x)$ such that $conf'(p_x, n) < Th$, we allocate k new workers w_1, \dots, w_k to record r_n for phase p_x , i.e. $W'_{in}(p_x, n) = W_{in}(p_x, n) \cup \{w_1, \dots, w_k\}$. The number of workers depend on the chosen policy (see details in Section 4). Accordingly, for every new worker w_i affected to a tagging task for a record r_n in phase p_x , we set $W'_{out}(p_x, n, i) = \emptyset$
- **Budget update.** We then update the budget. The overall number of workers hired is

$$nw = \sum_{p_x \in \mathcal{P}r_n} \sum_{r_n \in \mathcal{D}'_{in}(p_x)} |W'_{in}(p_x, n) \setminus W_{in}(p_x, n)|$$

We consider, for simplicity, that all workers and tasks have identical costs, we hence set $B' = B - nw$.

An execution begins from an initial configuration C_0 in which only p_0 is active, with an input dataset affected to p_0 , and starts with workers allocation. Executions end in a configuration C_f where all records in $\mathcal{D}_{in}(p_f)$ are tagged with a sufficient threshold. Notice that several factors influence the overall execution of a workflow. First of all, the way workers answers are aggregated influence the number of workers that must be hired to achieve a decent confidence in the synthesized answer. We propose to consider two main aggregation policies. The first one is majority voting (MV), where a fixed static number of workers is hired for each record in each phase. A second policy is the expectation maximization (EM) based technique proposed in [28], in which workers are hired on demand to increase confidence in the aggregated answer. With this policy, the confidence level is computed taking into account the estimated expertise of workers, and the difficulty of record tagging. The number of workers hired per record in a phase is not fixed, but rather computed considering the difficulty of tagging records, and the remaining budget.

For a given workflow, asynchrony is another key factor that may influence the time and budget spent to realize a complex task. Recall that for a phase p_x which input dataset is built as an union of sets of records, asynchronous guards allow to start processing records as soon as $\mathcal{D}_{in}(p_x) \neq \emptyset$. Synchronous guards force p_x to wait for the termination of its predecessors. In Section 5, we study the impact of synchronous/asynchronous guards on the overall execution of a workflow.

3 AGGREGATION MODEL

As mentioned in previous section, crowdsourcing requires replication of micro-tasks, and aggregation mechanisms for the answers returned by the crowd. For simplicity, we consider Boolean tasks, i.e. with answer 0 or 1. However, the model easily extends to a more general setting with a discrete set of answers.

Consider a phase p_x which input is a set of records $D_x = \{r_1, r_2, \dots, r_n\}$, and which goal is to associate a Boolean tag to each record of D_x . We assume a set of k independent workers that return Boolean answers, and denote by l_{ij} the answer returned by worker j for a record r_i . $L_i = \bigcup_{j \in 1..k} l_{ij}$ denotes the set of answers returned by k workers for a record r_i and $L = \bigcup_{j \in 1..n} L_j$ denotes the

set of all answers. We assume that workers are independent (there is no collaboration and their answers are hence independent), and *faithful* (they do not give wrong answers intentionally). The objective of aggregation is to derive a set of *final answers* $Y = \{y_j, 1 \leq j \leq n\}$ from the set of answers L . Once a final answer y_j is computed, it can be appended as a new field to record r_j .

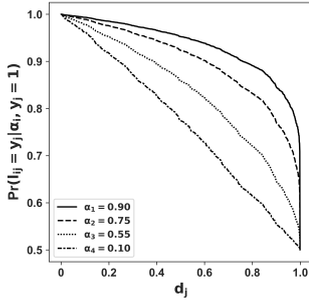


Figure 2: Generating functions $Pr(l_{ij} = y_j | d_j, \alpha_i, y_j = 1)$

The set of produced results can be used to launch new phases and to forward records to successor phases of p_x .

We consider several parameters to model tasks and workers, namely the difficulty to tag a record, and the expertise of workers. The *difficulty* to tag a record r_j is modeled by a real valued parameter $d_j \in [0, 1]$. Value 0 means that tagging r_j is very easy, and $d_j = 1$ means that it is extremely difficult. Expertise of a worker is often quantified in terms of *accuracy*, i.e. as the ratio of correct answers. However, accuracy can lead to bias in the case of datasets with unbalanced ground truth. Indeed, consider a case where the number of records with ground truth 1 is much higher than the number of records with ground truth 0. If a worker annotates most of records with ground truth 1 as 1 but makes errors when tagging records with ground truth 0, her accuracy will still be very high. We hence prefer a more precise model, where expertise of a worker is given as a pair $\xi_i = \{\alpha_i, \beta_i\}$, where α_i is the **recall** and β_i the **specificity** of worker i . The *recall* α_i is the probability that worker i answers 1 when the ground truth is 1, i.e. $\alpha_i = Pr(l_{ij} = 1 | y_j = 1)$. The *specificity* β_i is the probability that worker i answers 0 when the ground truth is 0, i.e. $\beta_i = Pr(l_{ij} = 0 | y_j = 0)$. We do not have a priori knowledge of the behavior of workers, so we define a generative model to determine the probability of correct answers when α_i, β_i are known. This probability depends on the difficulty of a task, on recall and specificity of the considered worker, and on the ground truth. We set $Pr(l_{ij} = y_j | d_j, \alpha_i, y_j = 1) = (1 + (1 - d_j)^{(1-\alpha_i)})/2$ and $Pr(l_{ij} = y_j | d_j, \beta_i, y_j = 0) = (1 + (1 - d_j)^{(1-\beta_i)})/2$.

Figure 2 shows probability to get $l_{ij} = 1$ when $y_j = 1$. The horizontal axis represents the difficulty of a task, the vertical axis denotes the probability to get answer $l_{ij} = 1$. Each curve represents this probability for a particular value of *recall*. Note that the vertical axis ranges from 0.5 to 1.0 as a random guess by a worker can still provide a correct answer with probability 0.5. As the difficulty of task increases, the probability of giving a correct answer decreases and when the task difficulty is 1 all workers cannot do better than a random guess. For a fixed difficulty of a task, the higher recall is, the more accurate answers are.

We equip complex workflows with an aggregation technique that uses Expectation Maximization (EM) [13] to estimate jointly latent variables $(\alpha_i)_{i \in 1..k}, (\beta_i)_{i \in 1..k}, (d_j)_{j \in 1..n}$ and derive the *final answer* y_j for each record r_j . We denote by θ the values of $(\alpha_i)_{i \in 1..k}, (\beta_i)_{i \in 1..k}, (d_j)_{j \in 1..n}$. EM iterates two alternating steps. In the E-step, we compute the posterior probability of $y_j \in \{0, 1\}$ for a given record r_j given the difficulty d_j , workers expertise $(\alpha_i, \beta_i)_{(i \in 1..k)}$ and the answers $L_j = \{l_{ij} \mid i \in 1..k\}$. In the M-Step, we compute the parameters θ that maximize $Q(\theta, \theta^t)$ with respect to the estimated posterior probabilities of Y computed during the E-step of the algorithm. Let θ^t be the value of parameters computed at step t of the algorithm. We use the observed values of L , and the previous expectation for Y . We maximize

$Q'(\theta, \theta^t) = \mathbb{E}[\log Pr(L, Y \mid \theta) \mid L, \theta^t]$ (we refer interested readers to [11]-Chap. 9 and [9] for explanations showing why this is equivalent to maximizing $Q(\theta, \theta^t)$). We can hence compute the next value as: $\theta^{t+1} = \arg \max_{\theta} Q'(\theta, \theta^t)$. We maximize $Q'(\theta, \theta^t)$

using optimization techniques provided by the `scipy`² library. We iterate E and M steps, computing at each iteration t the posterior probability and the parameters θ^t that maximize $Q'(\theta, \theta^t)$. The algorithm converges, and stops when the difference between two successive joint log-likelihood values is below a threshold (fixed in our case to $1e^{-7}$). It returns values for parameters $(\alpha_i)_{i \in 1..k}, (\beta_i)_{i \in 1..k}, (d_j)_{j \in 1..n}$. The final answer is the most probable y_j .

4 COST MODEL FOR WORKFLOW

The objective of a complex workflow W over a set of phases $P = \{p_0, \dots, p_x\}$ is to transform a dataset input to the initial phase p_0 and eventually produce an output dataset. The final answer is the result of the last processed phase p_f . The simplest scenario is a workflow that adds several binary tags to input records. The realization of a micro-task by a worker is paid, and workflows come with a fixed maximal budget B_0 provided by the client. For simplicity, we consider that each worker receives one unit of credit per realized task. As explained in section 2, each phase receives records, each record is tagged by one or several workers. Answers are then aggregated, and the records produced by a phase p_x are distributed to its successors if they meet some conditions on the data. A consequence of this filtering done by conditions is that records have different lifetimes and follow different paths in the workflow. Further, one can hire more workers to increase confidence in an aggregated result if needed and if a sufficient budget remains available. Several factors influence the realization of a workflow and its cost: the number of tagging tasks that have to be realized, the available initial budget, the confidence in produced results, workers expertise, the size and nature of input data, the difficulty of tagging, and the policies chosen to realize a workflow and to hire workers. Existing crowdsourcing platforms usually follow static allocation schemes, i.e. fix a number K_s of workers to hire for each micro-task. An obvious drawback of this approach is that the same effort is spent to solve easy and difficult tasks.

In section 2, we have defined synchronous and asynchronous schemes to allocate workers on-the-fly to tasks. In this section, we define the cost model associated with these schemes, and in particular, the threshold measure used to decide whether more workers should be hired. We show in section 5 that the algorithm achieves a good trade-off between cost and accuracy. Recall that at each round, we allocate new micro-tagging tasks to workers, to obtain answers for records that are still open. EM aggregation is used to obtain a plausible aggregated tag y_j^x for each record r_j from a set of answers L_j^x obtained in each active phase p_x . The algorithm gives an estimation of difficulty d_j^x of tagging record $r_j \in p_x$, and evaluates the expertise level of every worker w_i , i.e. its recall α_i and its specificity β_i . We also obtain a *confidence score* \hat{c}_j^x for the aggregated answer. This score is used to decide whether we need more answers or conversely consider y_j^x as a definitive result. Let $k_j^x = |L_j^x|$ denote the number of answers for record $r_j \in p_x$. The *confidence* \hat{c}_j^x in final label y_j^x is defined as:

$$\hat{c}_j^x (y_j^x=1) = \frac{1}{k_j^x} \cdot \sum_{i=1}^{k_j^x} \left\{ l_{ij}^x \times \left(\frac{1+(1-d_j^x)^{(1-\alpha_i)}}{2} \right) + (1 - l_{ij}^x) \times \left(1 - \frac{1+(1-d_j^x)^{(1-\alpha_i)}}{2} \right) \right\}$$

²docs.scipy.org/doc/scipy/reference/generated/scipy.optimize.minimize.html

$$\hat{c}_j^x(y_j^x=0) = \frac{1}{k_j^x} \cdot \sum_{i=1}^{k_j^x} \left\{ (1 - l_{ij}^x) \times \left(\frac{1+(1-d_j^x)^{(1-\beta_i)}}{2} \right) + (l_{ij}^x) \times \left(1 - \frac{1+(1-d_j^x)^{(1-\beta_i)}}{2} \right) \right\}$$

Confidence \hat{c}_j^x is a weighted sum of individual confidence of workers in the aggregated result. Each worker adds its probability of answering correctly (i.e., choose $l_{ij}^x = y_j^x$) when aggregating the final answer. This probability depends on y_j^x , but also on worker's competences. If this value is greater than a current threshold Th , then answer y_j^x is considered as definitive and the record r_j is closed. Otherwise, the record remains active. We fix a maximal number $\tau \geq 1$ of workers that can be hired during a round for a particular record. Let T_{ar} denote the set of active records after aggregation and D_{max}^x the maximal difficulty for a record of phase p_x in T_{ar} . For every record $r_j^x \in T_{ar}$ with difficulty d_j^x , we allocate $a_j^x = \lceil (d_j^x / D_{max}^x) \times \tau \rceil$ new workers for the next round. Intuitively, we allocate more workers to more difficult tasks. Now, T_{ar} and hence a_j^x depend on the threshold computed at each round. An appropriate threshold must consider the remaining budget, the remaining work to do, that depends on the number of records to be processed, on the structure of the workflow, and on the chosen policy. A first parameter to fix for the realization of a workflow is the initial budget B_0 .

Definition 4.1 (Height, width). The *height* of phase $p_x \in \mathcal{P}$ is the length of the longest path from the phase p_x to the final aggregated phase p_f and is denoted by h_x . The height a workflow W is the height of its initial phase, i.e. h_0 . The *width* of W is the size of the largest clique in W (i.e the largest subset $X \subseteq \mathcal{P}$ such that $\forall x, y \in X, x \rightarrow y$).

The notions of height and width are interesting to evaluate the required budget to terminate an execution of a workflow, and hence fix an appropriate threshold for confidence. The height of a workflow represents the minimal number of rounds needed to move a record from p_x to p_f . The width of a workflow is a bound on maximal number of active copies or a record at a given instant. Before considering how threshold is computed, we need to verify that a workflow has a sufficient budget for its execution. In each phase, a record is allocated at least $\tau \geq 1$ workers. So, if n is the total number of records to process in a workflow W , the budget spent will lay between $\tau.n.h_0$ and B_0 . Similarly, some records are sent to more than one successor, so in a workflow of width w , we may have to fund up to $\tau.n.h_0.w$ micro-tasks to terminate, but this is a very coarse approximation of the minimal resources needed to terminate a workflow. To obtain sharper evaluations, we first compute a bound on the number of remaining phases that records have to go through when they are currently processed in a phase p_x before completion of the workflow. We call this number the *foreseeable work* at phase p_x and denote it by $fw(p_x)$. We assume simple non-hierarchical structures, i.e. without nesting of exclusive and non-exclusive forks. For these structures, this bound can be computed iteratively according to the structure of the workflow³. Let p_x be a phase with several successors. If p_x is a non-exclusive fork phase, then there is no other fork along a branch before a merge phase. A phase p_y that is a merge phase immediately after p_x is called the corresponding phase of p_x . Intuitively, in a realization of W , all phases on a path between p_x and p_y can only process records that went through p_x . Let p_x be an exclusive fork phase. Then, the corresponding union node of

³A similar bound can be computed inductively for more complex workflows with nested exclusive/non-exclusive forks. We keep workflows simple for the sake of readability.

p_x is a node p_y such that all path originating from p_x visit p_y , and no predecessor of p_y satisfy this property. To decide whether p_y is the corresponding node of p_x one can compute the set of nodes that are both successors of p_x and predecessor of p_y , and decide for each of them whether they have a successor that is not a predecessor of p_y . This can be done in polynomial time. Then, the maximal number of phases that have to be realized between p_x and its corresponding node p_y is the length of the maximal path from p_x to p_y (which can also be computed in PTIME).

The algorithm to compute the foreseeable workload in a phase p_x consists in computing the most expensive path in a workflow once the cost of parallel processes contained between a non-exclusive fork node and its corresponding phase is evaluated and this set of nodes is replaced by a single phase of corresponding cost. This is described more precisely in Algorithm 1.

Algorithm 1: $FW(p_n)$

Data: Workflow W , phase p_n
Result: $fw(p_n)$

- 1 **for** each node p_x in $Succ(p_n)$ **do**
- 2 $Cost(p_x) := 1$;
- 3 **end**
- 4 **for** each non-exclusive fork node p_x in $Succ(p_n)$ **do**
- 5 Find p_y the corresponding node of p_x ;
- 6 $Z = Succ(p_x) \cap pre(p_y)$;
- 7 Replace Z by a fresh node p'_x ;
- 8 $Cost(p'_x) = |Z|$;
- 9 **end**
- 10 Compute the most expensive path $p_n.p_{i_1} \dots p_{i_k}.p_f$ from p_n to p_f ;
- 11 $fw := 1 + \sum Cost(p_{i_k})$;
- 12 **return** fw ;

Definition 4.2 (Foreseeable workload). Let n_x denote the total number of active records at a phase p_x in a configuration C . The *foreseeable task number* from p_x in configuration C is denoted $ft_C(p_x)$ and defined as $ft_C(p_x) = n_x \times FW(p_x)$. The *foreseeable task number* in configuration C is the sum $FTN(C) = \sum_{p_x \in \mathcal{P}} ft_C(p_x)$

Let us now define a threshold function based on the current configuration of the workflow. This function must consider all records that still need processing, the remaining budget, and an upper bound on the number of tagging tasks that will have to be realized to complete the workflow. Further, the execution policy will influence the way workers are hired, and hence the budget spent. In a synchronous execution, records in a phase p_x can be processed only when all records in preceding phases have been processed. On the contrary, in asynchronous execution mode, processing of records input to a phase p_x can start without waiting for closure of all records input to preceding phases. A consequence is that in synchronous modes, decisions can be taken *locally* to each phase, while in an asynchronous mode, the way to tune threshold must be taken according to a *global* view of the remaining work in the workflow. Hence, for an asynchronous execution policy, we will consider a global threshold function, computed for the whole workflow. On the contrary, for an asynchronous execution policy, we will define a more local threshold function computed phase by phase.

Asynchronous execution. We now define a global ratio $Rt \in [0, 1]$ of executed work in a workflow. The execution of a workflow starts from a configuration C_0 with an expected workload of $FTN(C_0)$. It is an upper bound, as all records do not necessarily visit the maximal number of phases. We define a ratio depicting the proportion of already executed or avoided work as $Rt = \frac{FTN(C_0) - FTN(C)}{FTN(C_0)}$. Note that at the beginning of an execution, $Rt = 0$ as no record is processed yet. When records are processed and moved to successor phases, Rt increases, and we necessarily have $Rt = 1.0$ when no record remains to process. Now, the threshold value has to account for the remaining budget to force the progress of records processing. Let B_{in} denote the budget at the beginning of execution, and B_c be the budget consumed. We denote by \mathcal{B} the fraction of budget consumed at a given execution time, i.e. $\mathcal{B} = \frac{B_c}{B_0}$. Note that, at the beginning of an execution, $B_c = \mathcal{B} = 0$. \mathcal{B} increases at every round of the execution, and takes value $\mathcal{B} = 1$ when the whole budget is spent. We now define a global threshold value $Th \in [0.5, 1.0]$ that accounts for the remaining work and budget.

$$Th = \frac{1 + (1 - \mathcal{B})^{Rt}}{2} \quad (1)$$

We remind that in a phase p_x , a record with confidence level $c > Th$ is considered as processed for phase p_x . In an asynchronous execution policy, the threshold is a global value and applies to all records in the workflow at a given instant. The intuition for Th is simple: when only a few record remain to be processed, and the remaining budget is sufficiently high, then one can afford final answers with a high confidence threshold. This means that many records will obtain new answers and probably increase their current confidence level. Conversely, if the number of records to be processed is high and the remaining budget is low, then the threshold decreases, and even records which current answer have a low confidence level are considered as processed and moved to the next phase(s).

Synchronous execution. In asynchronous execution, records are processed individually, and a phase does not wait for completion of its predecessors to start. As a consequence, all phases of the workflow can be active at the same time. As in each round, records can be processed in all phases, we consider a global threshold, and hence a global budget allocation policy. However, in synchronous executions, records are processed phase wise, i.e. a phase does not start processing its input dataset until all records in the preceding phases are processed. Here, the global threshold used in asynchronous executions may not work for synchronous executions. As task are realized phase wise, in the early phases of workflows execution, the threshold will be high. One may face situations where at the end of an execution, the number of records to process is still high, but a significant part of the budget has already been used. Within this setting, at the end of the execution, the threshold is low. It forces to accept final answers with low confidence. This may affect the overall quality of the final output of the workflow. To avoid this problem, we propose to allocate the budget phase by phase. The idea is to divide the budget among phases based on the number of records processed.

In synchronous executions, phases start processing all their records once preceding phases have completed their work (i.e. all records are processed and for each of them, a final answer with sufficient confidence score have been aggregated). We will say that a task becomes *active* when it starts processing records, and

denote by $Init(p_x)$ the number of records input to p_x when the phase becomes active.

As for asynchronous execution, synchronous execution starts with an initial budget B_0 , and in each configuration C , one knows the remaining budget $B_r(C)$. The key idea in synchronous execution is to compute resources needed for each active phase, a ratio of input records that still need additional answers to forge a trusted answer, and a local threshold.

The initial budget allocated to a phase p_x with n_x records in a configuration C is

$$B_0^x = \frac{B_r(C)}{\sum_{p_i \text{ active phase}} FTN(p_i)} \times n_x$$

Intuitively, one shares the remaining budget among active phases to allow termination of the workflow from each phase. Then for each phase, we maintain the consumed budget B_c^x for phase x , and the ratio $\mathcal{B}_x = \frac{B_c^x}{B_0^x}$ of consumed budget in phase p_x . Now, for each active phase p_x , we compute the ratio

$$R_t^x = \frac{|\{r_i \mid \hat{c}_i \leq Th_x\}|}{Init(p_x)} \quad (2)$$

where Th_x is the threshold computed for the previous round. A local threshold for the realization of the next round of an active phase can then be computed as in asynchronous execution, using the formula

$$Th_x = \frac{1 + (1 - \mathcal{B}_x)^{R_t^x}}{2} \quad (3)$$

With the convention that the initial threshold Th_x for a starting active phase, as no record is processed yet is $Th_x = \frac{1 + (1 - \mathcal{B}_x)}{2}$.

Realization of Workflows. Regardless of the chosen policy, the execution of a workflow always follows the same principles. The structure of workflow W is static and does not change with time. It describes a set of phases $P = \{p_0, \dots, p_f\}$, their dependencies, and guarded data flows from one phase to the next one. A set of n records $R = \{r_1, \dots, r_n\}$ is used as input to W , i.e., is passed to initial phase p_0 , and must be processed with a budget smaller than a given initial budget B_0 . As no information about the difficulty of a task d_j^x is available at the beginning of phase p_0 , τ workers are allocated to each record for an initial estimation round. The same principle is followed for each record when it enters a new phase $p_x \in W$. After collection of τ answers, at each round we first apply EM aggregation to estimate the difficulty d_j^x of active records $r_j \in p_x$, \hat{c}_j^x the confidence in the final aggregated answer y_j^x and the recall α_i and specificity β_i of each worker w_i . Then we use a stopping threshold to decide whether we need more answers for each of the records in a phase. In asynchronous execution, the threshold Th is a global threshold, and in synchronous mode, the confidence of each record r_j in a phase p_x is compared to the local threshold Th_x . Records with sufficient confidence are passed to the next phase(s), for other records we hire new workers to obtain more answers. This can increase the confidence level, but also decrease the threshold, as a part of the remaining budget is consumed. The execution stops when the whole budget B_0 is exhausted or when there is no additional record left to process. At the end, the final phase p_f returns the aggregated answer for each of the record.

Termination. One can easily see that as the remaining available budget decreases, the threshold used to decide whether the aggregated answer for a record is final decreases too. However, there are situations where the confidence in each answer remains

low, and the remaining budget reaches 0 before the threshold attains the lower bound 0.5 (that forces moving any record to the next phase(s)). Similarly, when records do not progress in the workflow, the ratio of remaining work R_t remains unchanged for many rounds. As a consequence, the realization of a workflow with our synchronous and asynchronous realization policies may not terminate. We will see in the experimental results section that even with poor accuracy of workers, this situation was never met. Non-termination corresponds to situations where the weighted answers of workers remain balanced for a long time. The threshold decreases slowly, and the confidence on aggregated answers remains lower. In that case, when threshold and confidence values coincide (in the worst case at value 0.5), the remaining budget is too low to realize the remaining work. Solutions to solve this issue and guarantee termination is to bound the sojourn time of a record in a phase, or to keep a sufficient budget to terminate the workflow with a static worker allocation policy hiring only a small number of workers per record.

5 EXPERIMENTS AND RESULTS

In this section, we evaluate execution policies on typical workflows. We consider a standard situation, where a client wants to realize a complex task defined by a workflow on a crowdsourcing platform. The client provides input data, and has a budget B_0 . We assume that crowd workers do not collaborate and hence realize their micro-tasks independently. As there exists no platform to realize complex task, there is no available data to compare the realization of a workflow with our approach to existing complex task executions. To address this issue, we design several typical workflows, synthetic data, and consider realizations of these workflows for various the execution policy, characteristics of data, and accuracy of workers.

We consider 5 different workflows, represented in Figure 4. Workflow W_1 is a sequence of tasks, W_2 is a standard fork-join pattern i.e. parallel processing of data followed by a merge of branches results, W_3 and W_4 are fork-join patterns with equal and different lengths on branches, and W_5 is a more complex workflow with two consecutive forks followed by merges on each branch. We consider that each micro task simply tags records, and simple exclusive guards sending each record to one successor, depending on the tag obtained at this phase. In Figure 4 we depict these choices by pairs of letters (l_0, l_1) representing the binary decision taken on each phase, and assume guards of the form $f == l_0$ or $f == l_1$, where f is the field of records produced by the phase. For example, in workflow W_1 , phase p_0 considers two possible tags denoted A and B . After realization of the tasks, if the records are tagged as A by the workers then records are moved to the phase p_f and if tagged with B the records are assigned to phase p_1 for further processing. Each phase of workflows implements similar tagging and decision.

We evaluate average costs and accuracies achieved by workflows realizations with the following parameters. First, the input of each complex task is a dataset of 80 records. Notice that despite this fixed size, the number of micro-tasks to realize by workers vary depending on the execution policy, on the value of data fields produced by workers, but also on the initial dataset, on the initial budget, etc. Each record in the original dataset has initially known data fields, and new fields are added by aggregation of workers answers during the execution of the workflow. For these fields, we assume a prior ground truth, which influences the probability that a worker answers 0 or 1 when filling this

| Workflow | W_1 | W_2 | W_3 | W_4 | W_5 |
|---------------------------------------------------------|-----------|-------------|--------------|-------|-------|
| Parameter | Value | | | | |
| Worker Accuracy | Low | Mid | Average | High | |
| Budget for B_{mv}, $k =$ | 10 | 20 | 30 | | |
| Data Type | Balanced | Imbalanced | | | |
| Mechanisms | Static MV | Synchronous | Asynchronous | | |

Table 1: Evaluation Parameters

field. We generate balanced (equal numbers of 0 and 1 in fields) and unbalanced datasets (unbalanced numbers of 0 and 1).

We run the experiment with 4 randomly generated pools of 50 crowd workers, making their accuracy range from low to high expertise. For each pool, we sampled accuracies of workers according to normal distributions ranging respectively in intervals $[0.2, 0.7]$ (low expertise of workers), $[0.4, 0.9]$ (low to average expertise), $[0.6, 0.99]$ (average expertise) and $[0.8, 0.99]$ (high expertise). The composition of pools is shown in Figure 3.

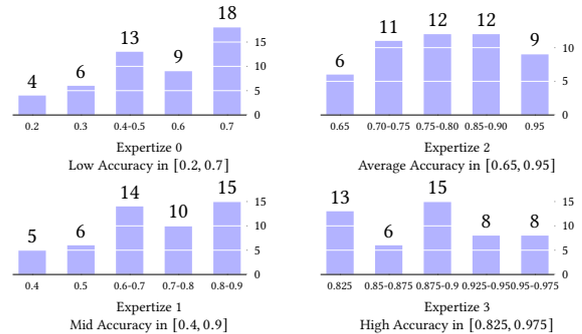


Figure 3: Distributions of workers accuracy.

The last parameter to set is the initial budget B_0 . We first evaluated the cost for the realization of workflows with a static allocation policy that associates a fixed number of k workers to each record in each phase, and aggregate their answers with Majority Voting. We call this policy *Static Majority Voting (SMV)*. For each workflow, we performed random runs of SMV to evaluate the maximal budget B_{mv} needed for three different values $k = 10/20/30$. Note that the total budget B_{mv} consumed by *staticMV* technique cannot be fixed a priori, as it depends on the execution path followed by records during execution, with random answers of workers. However, SMV is a naive approach, that was shown inefficient in most benchmarks [28], so starting with a budget $B_0 = B_{mv}$ for realization techniques tailored to save budget when accuracy is sufficient is a sensible approach.

In a second step, we used the total budget B_{mv} spent by the SMV approach as initial budget for synchronous and asynchronous execution. The idea is to achieve at least the same accuracy as SMV with synchronous and asynchronous execution with the same initial budget $B_0 = B_{mv}$, while spending a smaller fraction of this budget. Overall, our experiments cover realization of 5 different workflows with different values for initial budget, workers accuracy, characteristics of data, and realization policy. This represents 72 different contexts, represented Table 1 (one type of experiment represents a selection of one entry in each row). We run each experiment 15 times to get rid of bias. This represents a sample of 1080 workflow realizations.

We can now analyze the outcomes of our experiments. A first interesting result is that all workflow executions terminated without exhausting their given initial budget, even with low competences of users. A second interesting result is that for all realization policies, and for all workflows, complex workflows realization ends with poor accuracy when expertise is low. Consider for instance the results of Figure 5. This Figure gives the

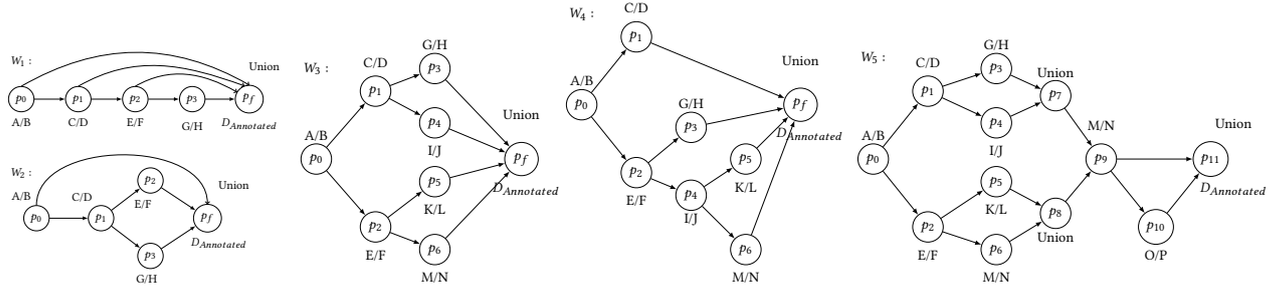


Figure 4: Five different workflows. W_1 : Sequence of phases, W_2 : Parallel data transformations followed by an aggregation of results, W_3 : Fork-join patterns with uniform lengths on their branches, W_4 : Fork-join patterns with nonuniform lengths on their branches, W_5 : Fork-join-Fork patterns with nonuniform lengths on their branches.

consumed budget and achieved accuracy for a given workflow and a given initial budget when workers have a low expertise. The first series of results concern Workflow 1 with a budget allowing respectively 10,20,30 workers per record in each phase. The overall expended budget with an SMV approach is around 1200, 2800, 4000, respectively. Regardless of the initial budget, synchronous and asynchronous approaches spend only a fraction of the budget allowed by SMV. Accuracy is not conclusive, as the best realization policy varies with each experiment: for instance, for W_1 with 10 workers per record to tag in each phase, SMV seems to be the best approach, while with a budget of 20, the synchronous approach is the best. However, most of the experiments achieve accuracies below 0.2, which is quite low. An explanation is that, as shown in Fig. 2, with low expertise, workers answers are almost random choices. Hence when all workers have a low expertise, individual errors are not corrected by other answers, and the ground truth does not influence the results. At each phase, the algorithms take their decisions mostly based on wrong answers provided by the low expert workers and in consequence the errors accumulate. The system’s behavior is then completely random, which results in poor performance. This tendency shown for all workflows and initial budgets with balanced data is confirmed on unbalanced data (the results of the experiments are shown in Appendix B, Figure 10).

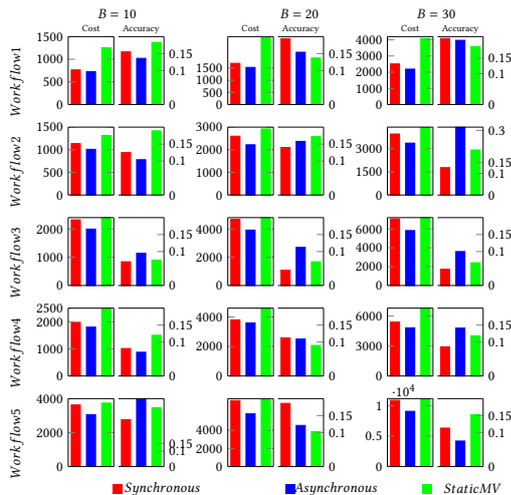


Figure 5: Budget and accuracy with low expertise

Next, we consider experiments with mid-level to high expertise, which is the most common case in a crowdsourcing setting. The experiments with competent workers and synchronous and

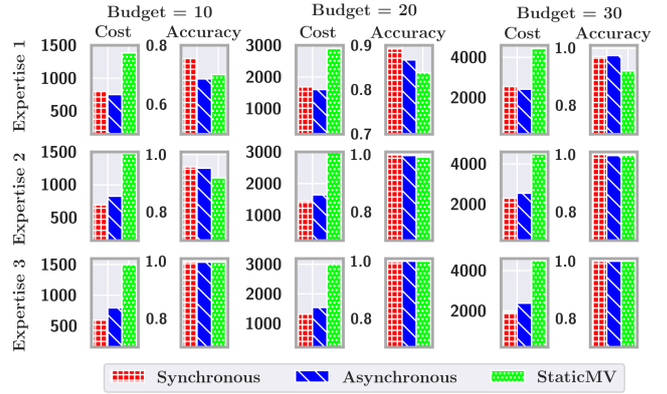


Figure 6: Workflow 1 on Balanced Data

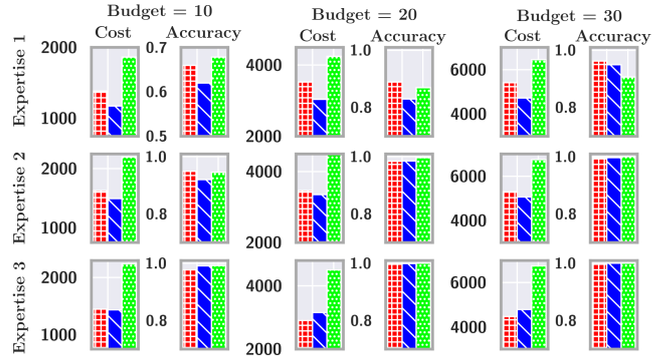


Figure 7: Workflow 1 on Unbalanced Data

asynchronous execution policies clearly show that dynamic allocation schemes outperform the *staticMV* approach both in terms of cost and accuracy. One can easily see these results Figures 6, 7, 8, and 9, that represent executions of workflows W_1 and W_2 with three levels of expertise, 3 initial budgets, and all execution policies, both for balanced and unbalanced data. We show similar results in Appendix B for workflows W_3 , W_4 , W_5 .

In the worst cases, *synchronous* and *asynchronous* executions achieve accuracies that are almost identical to that of *SMV*, but often give answers with better accuracy. With a sufficient initial budget, dynamic approaches achieve an accuracy greater than 0.9. An explanation for this improvement of synchronous and asynchronous executions w.r.t. *SMV* is that in *SMV*, one does not consider the expertise of the worker, whereas the *synchronous* and *asynchronous* executions are *EM* based algorithms that derive the final answers by weighting individual answers according to worker’s expertise. This makes *EM*-based evaluation of final answers more accurate than *static MV*. This improvement

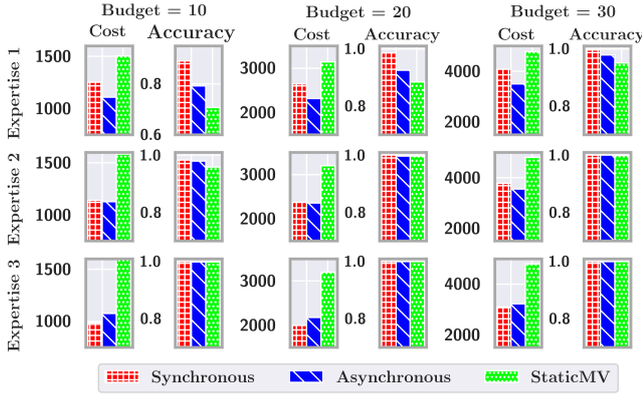


Figure 8: Workflow 2 on Balanced Data

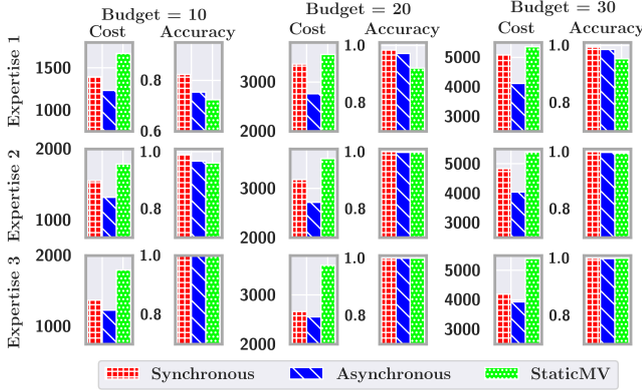


Figure 9: Workflow 2 on Unbalanced Data

already occurs at the level of a single phase execution (this was also the conclusion of [28]). The reasons for cost improvement with respect to *static MV* are also easy to figure. *SMV* allocates a fixed number of workers to every record in every phase of a workflow, whereas synchronous and asynchronous execution schemes allocate workers on-the-fly based on a confidence level which depends on the difficulty of tasks, workers expertise, and returned answers. By comparing confidence levels with a dynamic threshold, workers allocation considers the remaining budget and workload as well. This clever allocation of workers saves costs, as easy tasks call for the help of fewer workers than the fixed number imposed by *SMV*. The resources that are not used on easy tasks can be reused later for difficult tasks, hence improving accuracy.

These results were expected. A more surprising outcome of the experiment is that in most cases synchronous execution outperforms the asynchronous execution in terms of accuracy. The intuitive reason behind this result is that the way records are spread in the workflow execution affects the evaluation of expertise and difficulty. The synchronous execution realizes tasks in phases, while asynchronous execution starts tasks independently in the whole workflow. A consequence is that evaluation of hidden variables such as the difficulty of tasks and workers expertise evaluated by the EM aggregation improves with a larger number of records per phase in synchronous execution, while it might remain imprecise when the records are spread in different phases during an asynchronous execution. This precise estimation helps synchronous execution to allocate workers as well as to derive the final answers in a more efficient way and hence outperforms asynchronous execution. A third general observation is that both synchronous and asynchronous executions need a greater budget

to complete a workflow when data is unbalanced. Observe the results in figure 6 and Figure 7: the budgets spent are always greater with unbalanced data. A possible explanation is that in the balanced cases, records are distributed uniformly on all phases, which helps evaluation of workers expertise and difficulty of tasks, while with unbalanced data, some phases receive only a few records, which affects evaluation of hidden variables.

Unsurprisingly (see for instance Figure 6), for a fixed budget, when worker expertise increases, accuracy increases too, and consumed budget decreases. Competent workers return correct answers, reach a consensus earlier, and hence achieve better accuracy faster. Similarly for a fixed expertise level, increasing the initial budget increases the overall accuracy of the workflow. Again, the explanation is straightforward : a higher budget increases the threshold used to consider an aggregated answer as correct, giving better accuracies. To summarize, for a fixed initial budget and high enough expertise, synchronous and asynchronous policies usually improve both cost and accuracy.

6 CONCLUSION

In this work, we have proposed a framework to foster on the advantages of crowdsourcing systems and of workflow systems. The resulting model can be used to realize complex tasks with the help of a crowd of workers. A particular attention is paid to quality of the data produced, and to the overall cost of complex tasks realization. We have compared several task distribution strategies through experiments and showed that dynamic distribution of work outperforms static allocation in terms of cost and accuracy.

A short term extension for this work is to consider termination of complex tasks realization with dynamic policies. Indeed, workflows realized with dynamic policies may not terminate: this happens when, for some record, all workers agree to return the answers that do not increase the confidence. However, this situation was never met during our experiments, even with low expertise of workers. The probability of non-terminating executions with synchronous/asynchronous policies seems negligible. In our future work, we plan to demonstrate formally that $\mathbb{P}(B_r = 0 \wedge FTN(C) > 0)$, the probability of reaching a configuration with exhausted budget and remaining work to do is very low.

This work opens the way to new challenges. The next step is to test our approach with existing crowdsourcing platforms on a real case study. We are targeting citizen science initiatives, that typically require orchestration of various competence to reach a final objective. Now that our model is settled, another objective is to consider various strategies to hire workers in the most efficient way. A possibility to address this challenge is to see complex workflows as stochastic games, in which one player tries to maximize accuracy and reduce costs, while its opponent tries to achieve the opposite objectives.

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A APPENDIX: AGGREGATION WITH EM

$L_j = \{l_{1j}, \dots, l_{kj}\}$ denotes the answers (Observed labels) returned by k workers for a given task j . Here l_{ij} is the answer of worker i to task j . Expertise of k workers is modeled as recall $\alpha = \{\alpha_1, \dots, \alpha_k\}$ and specificity $\beta = \{\beta_1, \dots, \beta_k\}$. Given the observed answers of workers, objective is to infer the final label y_j , and

to derive the most probable values for d_j, α_i, β_i . We use a standard EM approach to infer the most probable actual answer $Y = \{y_1, \dots, y_n\}$ along with the latent variables $\Theta = \{d_j, \alpha_i, \beta_i\}$. E and M phases of the algorithm is illustrated below.

E Step: We assume that all the answers L are independently given by the workers. There is no collaboration between them. So, in every $L_j = \{l_{1j}, \dots, l_{kj}\}$, l_{ij} 's are independently sampled variables. We compute the posterior probability of $y_j \in \{0, 1\}$ for a given task j given the difficulty of task d_j , worker expertise $\alpha_i, \beta_i, i \leq k$ and the worker answers $L_j = \{l_{ij} \mid i \in 1..k\}$. Using Bayes' theorem, for a particular value $\lambda \in \{0, 1\}$ we have:

$$Pr[y_j = \lambda | L_j, \alpha, \beta, d_j] = \frac{Pr(L_j | y_j = \lambda, \alpha, \beta, d_j) \cdot Pr(y_j = \lambda | \alpha, \beta, d_j)}{Pr(L_j | \alpha, \beta, d_j)} \quad (4)$$

Observe that, that y_j and α_i, β_i, d_j are independent variables. We assume that both values of y_j are equiprobable, i.e. $Pr(y_j = 0) = Pr(y_j = 1) = \frac{1}{2}$. Hence we get:

$$Pr[y_j = \lambda | L_j, \alpha, \beta, d_j] = \frac{Pr(L_j | y_j = \lambda, \alpha, \beta, d_j) \cdot Pr(y_j = \lambda)}{Pr(L_j | \alpha, \beta, d_j)} = \frac{Pr(L_j | y_j = \lambda, \alpha, \beta, d_j) \cdot \frac{1}{2}}{Pr(L_j | \alpha, \beta, d_j)} \quad (5)$$

Similarly, the probability to obtain a particular set of labels L_j is given by:

$$Pr(L_j | \alpha, \beta, d_j) = \frac{1}{2} \cdot Pr(L_j | y_j = 0, \alpha, \beta, d_j) + \frac{1}{2} \cdot Pr(L_j | y_j = 1, \alpha, \beta, d_j) \quad (6)$$

Overall we obtain:

$$Pr[y_j = \lambda | L_j, \alpha, \beta, d_j] = \frac{Pr(L_j | y_j = \lambda, \alpha, \beta, d_j)}{Pr(L_j | y_j = 0, \alpha, \beta, d_j) + Pr(L_j | y_j = 1, \alpha, \beta, d_j)} \quad (7)$$

Let us consider one of these terms, and let us assume that every l_{ij} in L_j takes a value λ_s . We have

$$Pr(L_j | y_j = \lambda, \alpha, \beta, d_j) = \prod_{i=1}^k Pr(l_{ij} = \lambda_s | \alpha_i, \beta_i, d_j, y_j = \lambda) \quad (8)$$

If $\lambda_s = 0$ then $Pr(l_{ij} = \lambda_s | \alpha_i, \beta_i, d_j, y_j = 0)$ is the specificity, i.e. the probability to classify correctly a 0 as 0, and is the value $\delta_{ij} = \frac{1 + (1 - d_j)^{(1 - \beta_i)}}{2}$. Similarly, if $\lambda_s = 1$ then $Pr(l_{ij} = \lambda_s | \alpha_i, \beta_i, d_j, y_j = 1)$ is the recall, i.e. the probability to classify correctly a 1 as 1, and is the value $\gamma_{ij} = \frac{1 + (1 - d_j)^{(1 - \alpha_i)}}{2}$. Then the probability to classify $y_j = 1$ as $\lambda_s = 0$ is $(1 - \gamma_{ij})$ and the probability to classify $y_j = 1$ as $\lambda_s = 1$ is $(1 - \delta_{ij})$. We hence have $Pr(l_{ij} = \lambda_s | \alpha_i, \beta_i, d_j, y_j = 0) = (1 - \lambda_s) \cdot \delta_{ij} + \lambda_s \cdot (1 - \gamma_{ij})$. Similarly, we can write $Pr(l_{ij} = \lambda_s | \alpha_i, \beta_i, d_j, y_j = 1) = \lambda_s \cdot \gamma_{ij} + (1 - \lambda_s) \cdot (1 - \delta_{ij})$. So equation 7 rewrites as :

$$\begin{aligned} Pr[y_j = \lambda | L_j, \alpha, \beta, d_j] &= \frac{\prod_{i=1}^k Pr(l_{ij} = \lambda_s | y_j = \lambda_s, \alpha_i, \beta_i, d_j)}{Pr(L_j | y_j = 0, \alpha, \beta, d_j) + Pr(L_j | y_j = 1, \alpha, \beta, d_j)} \\ &= \frac{\prod_{i=1}^k (1 - \lambda_s) \cdot [(1 - \lambda_s) \delta_{ij} + \lambda_s (1 - \gamma_{ij})] + \lambda_s \cdot [\lambda_s \gamma_{ij} + (1 - \lambda_s) (1 - \delta_{ij})]}{Pr(L_j | y_j = 0, \alpha, \beta, d_j) + Pr(L_j | y_j = 1, \alpha, \beta, d_j)} \quad (9) \\ &= \frac{\prod_{i=1}^k (1 - \lambda_s) \cdot [(1 - \lambda_s) \delta_{ij} + \lambda_s (1 - \gamma_{ij})] + \lambda_s \cdot [\lambda_s \gamma_{ij} + (1 - \lambda_s) (1 - \delta_{ij})]}{\prod_{i=1}^k (1 - \lambda_s) \delta_{ij} + \lambda_s (1 - \gamma_{ij}) + \prod_{i=1}^k \lambda_s \gamma_{ij} + (1 - \lambda_s) (1 - \delta_{ij})} \end{aligned}$$

In the E step, as every α_i, β_i, d_j is fixed, one can compute $\mathbb{E}[y_j | L_j, \alpha_i, \beta_i, d_j]$ and also choose as final value for y_j the value $\lambda \in \{0, 1\}$ such that $Pr[y_j = \lambda | L_j, \alpha_i, \beta_i, d_j] > Pr[y_j = (1 - \lambda_s) | L_j, \alpha_i, \beta_i, d_j]$. We can also estimate the likelihood for the values of variables $P(L \cup Y | \theta)$ for parameters $\theta = \{\alpha, \beta, d\}$, as $Pr(y_j = \lambda, L | \theta) = Pr(y_j = \lambda_s, L) \cdot Pr(L_j | y_j = \lambda_s, \theta) = Pr(y_j = \lambda_s) \cdot Pr(L_j | y_j = \lambda_s, \theta)$

M Step: In M-Step, we compute the parameters θ that maximize $Q(\theta, \theta^t)$ with respect to the estimated posterior probabilities of Y computed during the E phase of the algorithm, Let θ^t be the value of parameters computed at step t of the algorithm. We use the observed values of L , and the previous expectation for Y . We maximize $Q'(\theta, \theta^t) = \mathbb{E}[\log Pr(L, Y | \theta) | L, \theta^t]$ (we refer interested readers to [11]-Chap. 9 and [9] for explanations showing why this is equivalent to maximizing $Q(\theta, \theta^t)$). We can hence

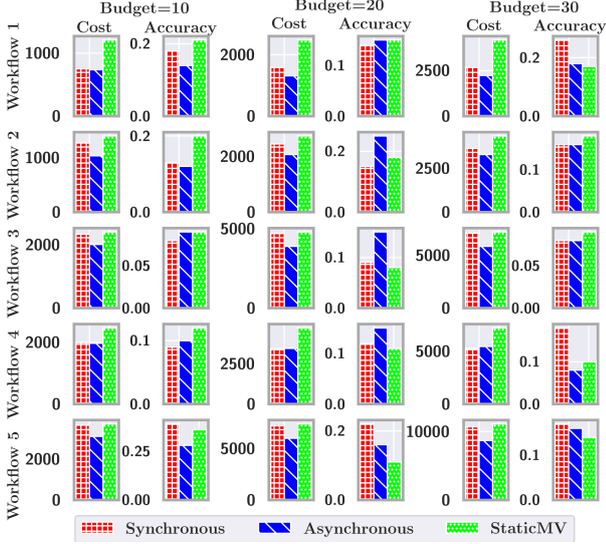


Figure 10: Results for low expertise & unbalanced Data

compute the next value as: $\theta^{t+1} = \arg \max_{\theta} Q'(\theta, \theta^t)$. Here in our context the values of θ are α_i, β_i, d_j . We maximize $Q'(\theta, \theta^t)$ using bounded optimization techniques provided by the standard SCIPY implementation. We iterate E and M steps, computing at each iteration t the posterior probability and the parameters θ^t that maximize $Q'(\theta, \theta^t)$. The algorithm converges, and stops when the improvement (difference between two successive joint log-likelihood values) is below a threshold (in our case $1e^{-7}$).

B ADDITIONAL RESULTS

For completeness, we give the experimental results for low accuracy (Fig. 10), and for the 72 contexts considered (all workflows, 3 initial budgets, all levels of expertise, and all types of data).

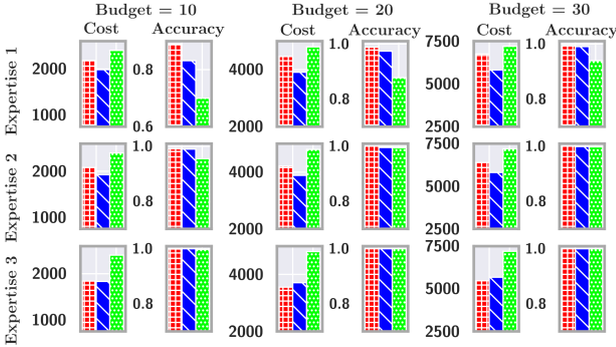


Figure 11: Workflow 3 on Balanced Data

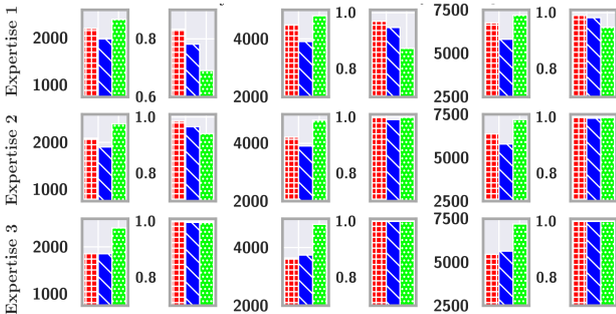


Figure 12: Workflow 3 on Unbalanced Data

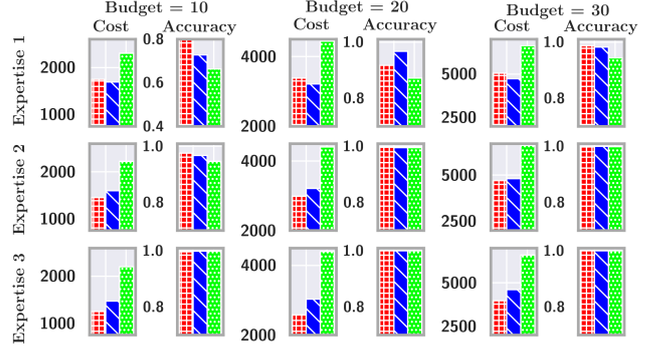


Figure 13: Workflow 4 on Balanced Data

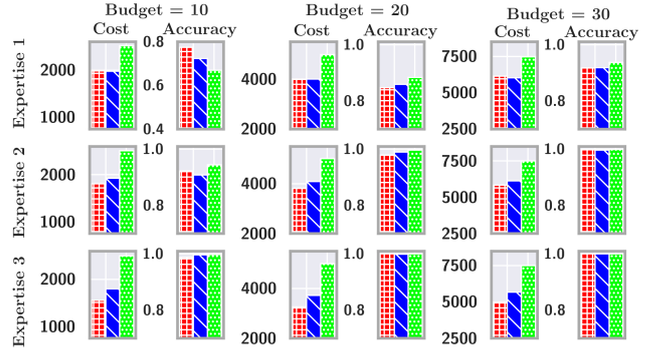


Figure 14: Workflow 4 on Unbalanced Data

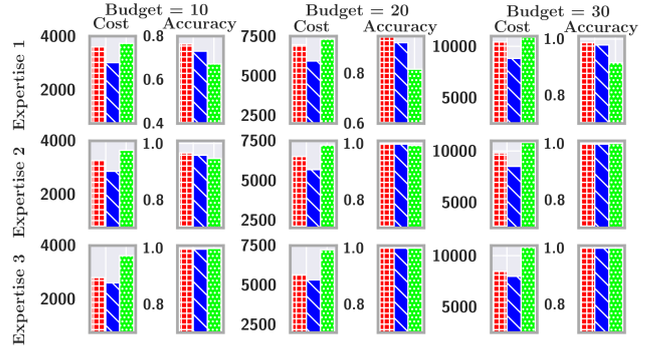


Figure 15: Workflow 5 on Balanced Data

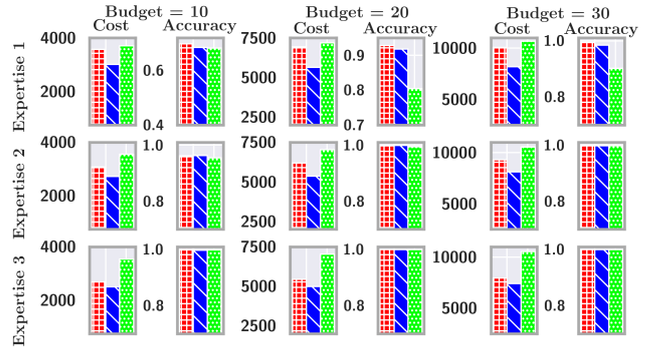


Figure 16: Workflow 5 on Unbalanced Data