**SLearn**: Shared learning human activity labels across multiple datasets

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**Abstract**—The research of sensor-based human activity recognition has been attracting increasing attention over years as it is playing an important role in various human-beneficiary applications such as ambient assistive living, health monitoring, and behaviour changing. Nowadays, the advancement of sensing and communication technologies has led to the possibility of collecting a large amount of sensor data, however, to build a reliable computational model and accurately recognise human activities we still need the annotations on sensor data. Acquiring high-quality, detailed, continuous annotations is a challenging task. In this paper, we explore the solution space on sharing annotated activities across different datasets in order to enhance the recognition accuracies. We have designed and developed two approaches: sharing training data and sharing classifiers towards addressing this challenge. We have validated the approach on three datasets and demonstrated their effectiveness in recognising activities only with annotations from as little as 0.1% of each dataset.

**I. Introduction**

Sensor-based activity recognition has increasingly played a significant role in many applications, which enables to provide customised services to suit people’s current context; for example, better optimising energy by adjusting heaters based on users’ activity routine [4], understanding the impact of human behaviours to environmental phenomena such as the change of indoor air quality [12], or assessing physical and cognitive conditions of disadvantaged residents [23].

Human activity recognition has been a popular research topic in the last decade, and numerous machine learning, data mining, knowledge-driven techniques have been applied and achieved promising results at targeting various problems. However, challenges still remain, and a crucial challenge resides in the scarcity of annotated datasets. First of all, acquiring annotations from users can be a time- and effort-consuming task. The state-of-the-art activity recognition technologies either use video to record activities of users, which will later be annotated by third parties, or rely on users’ self-report. The former approach has an undesirable privacy concern, while the latter can result in less accurate, incomplete annotations [26]; for example, users forget to record certain activities or use different terms to describe their activities.

Even though the recent attempts have helped to address the above challenges by facilitating self-annotation with the support of technologies [27] and enhancing the consistency of annotation labels via ontologies [19], acquiring annotations for long-term activity recognition is still a challenging task. That is, we only collect ground truth over the training period (e.g., the first few weeks or months) so that the machine learning techniques can use the annotated training data to build models. After that, we assume that the models stay fixed and can be used continuously to recognise human activities in the long term (e.g., for a couple of years). However, this assumption does not consider the fact that human activity routine or activity pattern might change over time, users (or background users like visitors or even pets) might interact with sensors differently, or users might conduct new activities that have not been recorded at all during the training period; that is, they might behave differently from the training period to the actual system running period.

Therefore, acquiring annotations to enable long-term activity recognition is one of the main challenges, and there have been different approaches proposed to address towards this problem. Unsupervised activity recognition is one way, where we either rely on common sense knowledge from the other sources (such as the Web or other textual sources [29] or domain knowledge [31]) to build activity profiles or build computationally expensive models to separate sensor data for each type of activities [22]. Another direction is active learning, where a classifier will build a computational model on a limited set of annotated data, and then iteratively annotate a large amount of un-annotated data. In each iteration, it will select uncertain data points that sit at the boundary of classes, and ask the human operators to annotate them so as to use them further train and refine the computational model. This approach can significantly reduce human annotation effort, however, it still requires all the activities of interest are covered in the initial annotated data. Recently transfer learning techniques are starting to attract attention and have also been applied to address this challenge. For example, a meta-feature space mapping technique has been developed to map sensors from one home environment to another so as to share the activity recognition model [5], but this approach can be subject to the density of sensors being deployed and the routines of activities from different users.

In this paper, we hypothesise that activity recognition accuracies can be improved by shared learning across multiple datasets to complement the insufficient amount of labelled data. That is, if each dataset contributes to a very limited number of labelled examples (even with an incomplete list of activities of interest), through shared learning between these
datasets, we are able to better recognise unlabelled examples than classic activity recognition approaches. The contribution of this paper is to propose algorithms for shared learning across activity recognition models from different users in different environments with different deployments of sensors. We evaluate the algorithms on three third-party datasets and demonstrate their effectiveness by comparing them against the state-of-the-art classification techniques.

The rest of the paper is organised as follows. Section II proposes the problem that SLearn aims to address with formal definitions and an illustration example. Section III reviews the mainstream work on addressing the scarcity of labelled data and identifies the difference between them and SLearn. Section IV describes the SLearn approach and Section V introduces the evaluation methodology and experiment setup. Section VI performs an evaluation and discusses the strength and limitation, and Section VII concludes with some suggested future work.

II. PROBLEM STATEMENT

In this section, we will introduce the research problem that the SLearn tries to address, through a formal definition and a concrete illustrative example.

A. Formal Definition

Based on the syntax and terminology from transfer learning [16], we first give preliminary definitions of problems that the SLearn approach aims to address.

Definition 1. Let a dataset DS consist of a domain D and a task T, where

- a domain D is a two-tuple \((\chi, P(X))\), \(\chi\) is the feature space of D and \(P(X)\) is the marginal distribution where \(X = [x_1, \ldots, x_n] \in \chi\).
- a task T is a two-tuple \((Y, f())\) for a given domain D. Y is the label space of D, which is a collection of possible labels \(Y = \{y_1, y_2, \ldots, y_m\}\). \(f()\) is an objective predictive function for D.
- Within this dataset, each pair \((X, y) \in \chi \times Y \cup \{\Theta\}\), where \(\Theta\) represents unknown or unlabelled. That is, some instances in the feature space have labels; i.e., \((X, y) \in \chi \times Y\) and \(f(X) \in Y\), while the other instances do not; i.e., \((X, \Theta) \in \chi \times \{\Theta\}\) and \(f(X) = \Theta\).

Definition 2. Let \(DSS\) be a collection of datasets \(\{DS_1, DS_2, \ldots, DS_l\}\), where each dataset has different domains (i.e., \(\forall 1 \leq i, j \leq L, i \neq j, \chi_i \neq \chi_j\) and \(P_i \neq P_j\)) and has different tasks (i.e., \(Y_i \neq Y_j\) and \(f_i \neq f_j\)). SLearn aims to assign a label to each unlabelled instance in the dataset; that is, \(\forall X \in \chi_i\), there exists a label \(y \in Y_1 \cup Y_2 \cup \ldots \cup Y_L\). To make it work, there are two assumptions:

- Feature space remapping - the feature spaces between these datasets are comparable and can be remapped between each other. That is, there exists a mapping function \(\theta_{i,j} : \chi_i \rightarrow \chi_j\) that remaps each instance \(X \in \chi_i\) to an instance \(X'\) in \(\chi_j\).
- Label space remapping - the label spaces between these datasets are comparable and can be remapped between each other. That is, there exists a mapping function \(\vartheta_{i,j} : Y_i \rightarrow Y_j\) that remaps each label \(y\) in \(Y_i\) to a label \(y'\) in \(Y_j\).

B. An Illustrative Example

Here we will illustrate the above definitions through an example of the SLearn scenario, which is activity recognition in a smart home environment deployed with binary event-driven sensors. Assume we deploy sensors in two different home settings: House I and House II. Due to different spatial layout and users’ preference, we deploy a different number of sensors with different sensing technologies in these two environments. For example, one house is deployed with a dozen of infra-red passive motion sensors, indicating the user’s whereabouts, while the other house is deployed with switch sensors to monitor the open/close states of cupboards, doors, or windows, and the pressure sensors to indicate whether a user sits on sofa or sleeps in bed.

When the sensors are deployed, we can start activating a SLearn Algorithm, without the need for explicit training period, as the training is continuously ongoing. Both users can start annotating their activities at their own pace. For example, if the user in House I has annotated a few examples of “preparing breakfast” and the user in House II has annotated a few examples of “taking bath”, then the SLearn Algorithm aims to recognise these two activities in both House I and II. It will support long-term incremental activity recognition in that it can accommodate new activities annotated by different users over time.

III. RELATED WORK

Activity recognition has been an active research topic in the last decade, and a large number of knowledge- and data-driven techniques have been proposed [30]. Among them, different approaches have been designed to address the scarcity challenge of activity annotation, including unsupervised learning, activity learning, and transfer learning. In the following, we will compare and contrast these approaches with SLearn.

A. Unsupervised Learning

Unsupervised learning automatically partitions and characterises sensor data into patterns that can be mapped to different activities without the need of annotated training data. Pattern mining and clustering are the two mostly used techniques that support unsupervised activity recognition. Gu et al. have applied emerging patterns to mine the sequential patterns for interleaved and concurrent activities [7]. Rashidi et al. propose a method to discover the activity patterns and then manually group them into activity definitions [18]. Based on the patterns, they create a boosted version of a Hidden Markov Model (HMM) to represent the activities and their variations in order to recognise activities in real time. Similarly, Ye et al. have combined the sequential mining and clustering algorithms to discover representative sensor events for activities. Different
from the work in [18], they have applied the generic ontologies to automatically map the discovered sensor sequential patterns to activity labels through a semantic matching process [31]. Yordanova et al. have also applied domain knowledge in rule-based systems to generate probabilistic models for activity recognition [11], [33].

Taking a different route, researchers also have applied web mining and information retrieval techniques to extract the common-sense knowledge between activities and objects via mining online documents; that is, what objects are used to perform a daily activity and how significant each object is contributed to identifying this activity [15], [29], [32], [34]. During the reasoning process, the mined objects are mapped to sensor events and an appropriate activity will be recognised.

SLearn is not a classic activity recognition problem where for each dataset we train a model with labelled instances and recognise unlabelled instances. In SLearn, the training data is assumed to be incomplete in that it might not cover all the activities of interest. SLearn is not a unsupervised learning technique as it still relies on labelled training data, but the labels can come from different datasets.

B. Active Learning

Active learning, so called “query learning”, is a subfield of machine learning, which is motivated by the scenario when there is a large amount of unlabelled data but a limited and insufficient amount of labelled data. As the labelling process is tedious, time-consuming and expensive in real-world applications, active learning methods are employed to alleviate the labelling effort by selecting the most informative instances to be annotated [20].

Alemdar et al. apply active learning strategies to select the most uncertain instances to be annotated; that is, the instances sit at the boundaries of different activity classes [1]. The annotated instances are then used to iteratively update a HMM to infer daily activities in a home setting. Their experimental results have demonstrated that active learning strategies have improved recognition accuracies, compared to random selection. Cheng et al. apply a density-weighted method that combines both uncertainty and density measure into an objective function to select the most representative instances for user annotation, which has been demonstrated to improve activity recognition accuracy with the minimal labelling effort [3]. Similarly, Hossain et al. combine the uncertainty measure and Silhouette coefficient to select the most informative instances as a way to discover new activities [8].

SLearn is not an active learning problem [20] in that it will not query a user or a human operator, but learns labels from the other datasets. However, we could use the uncertainty sampling strategies in active learning to determine when an algorithm should leverage training data or classifiers from other datasets.

C. Transfer Learning

Transfer learning is another approach to deal with the limitation of labelling data, where knowledge learned from a source domain (with labelled data) can be transferred to a target domain (without labelled data) [16].

Zheng et al. [34] propose an algorithm for cross-domain activity recognition that transfers the labelled data from a source domain to a target domain so that the activity model in the source domain can help to complete the similar activity model in the target domain. The similarity is not only measured on the objects being involved in the activities, but also on their underlying physical actions. One example is that the activity ‘Washing-laundry’ is similar to “Hand-washing dishes” on the action of “Hand washing”. They use the web search and apply the information retrieval techniques to build the similarity function that produces different probabilistic weights of actions and objects on activities of interest. These weights will be further used to train a multi-class weighted support vector machine to support activity recognition.

Maekawa et al. [13] have proposed an unsupervised approach to recognise physical activities from accelerometer data. They utilise information about users’ characteristics such as height and gender to compute the similarity between users, and find and adapt the models for the new users from the similar users.

van Kasteren et al. [24] propose a manual mapping between sensors in different households and learn the parameters of a target model using the EM algorithm to transit probabilities of HMM models from source to target. Similarly, Rashidi et al. [17] learn sensor mappings based on their locations and roles in activity models. The role is characterised in mutual information, measuring the mutual dependence between an activity and a sensor and suggests the relevance of using the sensor in predicting the corresponding activity. Feuz et al. [5] propose a data-driven approach to automatically map sensors based on their meta-features, which are mainly about when a sensor reports, and time intervals between events reported by this sensor and other sensors.

SLearn is most relevant to but not the same as transfer learning [16]. Our assumption is slightly different from the above works where they assume a complete model (that is, containing all the activities of interest) can be learnt on a source domain, while we assume each domain can only have a small fraction of data being annotated (that is, the activities having been annotated can be a subset of activities of interest in a domain) and we do not assume any domain necessarily as a source or target domain. However, transfer learning techniques such as feature remapping can be applied to SLearn. Especially, our approach is most similar to the above three, where we focus on sensor mappings to support sharing sensor data across multiple datasets. The difference is that we are using a knowledge-driven approach where sensors are modelled in location and object ontologies whose generality across different households and sensing technologies has been demonstrated in other works [31].

D. Co-training

The co-training algorithm, proposed by Blum and Mitchell in 1998, has been one classic approach to be able to boost
performance of a learning algorithm by leveraging a large number of unlabelled examples [2]. The idea is that the
description of each example can be partitioned into two distinct views, and each view can be linked with edges in a bipartite
graph. Then two classifiers can be trained separately on each view, and then results from each classifier are used to enlarge
the training set of the other. The co-training algorithm and its
variations have been recently applied in the multi-view human
activity recognition in smart home environments [6]. That is,
an activity can be viewed from different platforms of sensor
streams, such as acceleration data, motion sensor, or video.
The principle is to learn the same activities from each sensor
platform and share and adapt labels from one sensor platform
to another. For example, a home is equipped with ambient
sensors to monitor motion, lighting, temperature, and door
use. The resident now wants to train smart phone sensors to
recognise the same activities on ambient sensors. Whenever
the phone is located inside the home, both sensing platforms
collect data while activities are performed, resulting in a multi-
view learning opportunity where the ambient sensors represent
one view and the phone sensors represent the other view. If the
phone can be trained, it can also monitor activities outside of
the home and can update the home’s model when the resident
returns. The phone may converge upon a stronger model than
the home either because it receives training data of its own or
because it has a more expressive feature space.

The SLearn problem does not share the same assumption
as the co-training approach in that the co-training approach
works on multiple views from the same data while the
SLearn approaches need to work on multiple datasets that
share the sensing mission; that is, compatible sensor features
and activities.

IV. PROPOSED APPROACH

We hypothesise that we can significantly improve activity
recognition accuracies by leveraging a small set of training
data contributed from each dataset. All these datasets might
have a different number of sensors, in different sensing
technologies, deployed in different environments, and from
different users, as long as they have compatible feature space
and activities of interest. The contributed training data might
only contain a couple of examples, and possibly not cover the
whole set of activities of interest. In the following, we will
explore the solution space to address this question.

First of all, a classic approach of dealing with a small
amount of training data is leveraging unlabelled data, which
is similar to the active learning approach, mentioned in Sec-
tion III-B. That is, for each dataset, we train a classifier
on its labelled data and then use it to infer the labels on its unlabelled examples for $T$ rounds or until the
algorithm converges. For each iteration, we select the top $k$
most confident examples to enlarge the labelled data pool and
iteratively update the classifier. However, given our assumption
that the labelled data might be too little and have not covered
the whole set of activities of interest, this basic approach can
only assign the labels that have been observed in the training
data. We will need to leverage labels from the other datasets.
To do so we will look into two directions: sharing training
data and sharing classifiers.

A. Sharing training data

To share training data, we will need to map data from
the feature space of one dataset to the feature space of the
other dataset. An intuitive approach is feature-space remap-
ing in transfer learning; that is, convert an instance $X_I = \{x_{I,1}, x_{I,2}, ..., x_{I,n_I}\}$ from a dataset $I$ to a representation in an-
other dataset $\forall I, n_I$, $X_{II} = \{x_{II,1}, x_{II,2}, ..., x_{II,n_{II}}\}$. There are different feature remapping strategies introduced in [16]. Feuz et al. [5] have proposed a meta-feature based
mapping function for event-driven sensors in smart home
environments. They have defined a range of meta-features
about each sensor; for example, the average sensor event
frequency over 1-hour time periods, over 3/8/24-hour periods,
the mean and standard deviation of the time between this
sensor event and the next sensor event, and the probability of
the next event is from the same sensor. These meta-features
are used as a heuristic to guide the mapping process. This
is a data-driven approach for feature-space remapping, but
its performance might be affected by the activity routine
of various users and the deployment and types of sensing
technologies in each environment. For example, one user
might often have breakfast at 6am while the other might have
at 9am, or one user prefers having shower before breakfast
while the other prefers the other way around so that the
sensors might be mis-matched on the time scale. In addition,
the density of the sensor deployment and the frequencies and
sensitivity of sensors reporting events might affect the mapping
on the intervals between events. For example, one environment
can be more densely deployed with sensors so that the time
distances between events reported by different sensors can be
significantly shorter than the other set up with much fewer
sensors.

To reduce the impact of such differences in each dataset,
we will adopt a more general approach - semantics-based
feature mapping. We will use the common knowledge [31],
which has demonstrated generality across different smart home
datasets. The principle of semantics-based feature mapping is
to compute similarity between a pair of sensors based on where
they are deployed and which object they have attached to. Both
location and object concepts are organised in an ontological
hierarchy, from which a conceptual similarity measure [28] is
applied to calculate the distance between two concepts. Here
we re-apply the conceptual ontologies to feature re-mapping.

**Definition 3.** Given an instance $X_I = \{x_{I,1}, x_{I,2}, ..., x_{I,n_I}\}$, a semantics-based feature mapping function is defined as
\[
\theta(X_I) = X_{II}, \text{denoted as } \{x_{II,1}, x_{II,2}, ..., x_{II,n_{II}}\}, \text{where}\n\forall 1 \leq j \leq n_{II}, x_{II,j} = \sum_{i \in S_j} \frac{x_i \times \text{sim}(s_i, s_{II,j})}{|S_j|}, \text{where } S_j \text{is a collection of sensors in the feature space } I \text{ that are similar to the sensor } j \text{ in II.}
\]

\[
|S_j| \{s_i|1 \leq i \leq n_I, \text{sim}(s_i, s_{II,j}) > \varepsilon\}, \text{sim}(s_i, s_{II,j}) = w_L \times \text{sim}_L(s_i, s_{II,j}) + w_O \times \text{sim}_O(s_i, s_{II,j}), \text{where } w_L + w_O = 1.
\]
where $\varepsilon$ is the threshold to choose similar sensors, $sim_L$ and $sim_O$ are the similarity measure on location and object concepts that the $i$th sensor in I and $j$ sensor in II, and $w_L$ and $w_O$ are the weights on the location and object similarity.

Based on the above definition 3, a value $x_{iI,j}^{*}$ in a converted instance $X_{iI}^{*} \in \chi_{II}$ is the weighted average of the probabilities of all the similar sensors in the source feature space $\chi_I$ to the $j$th sensor in $\chi_{II}$ and the weight is their sensor similarity. That is, we try to estimate the probability of the $j$th sensor reporting events by looking at the probabilities of all of its similar sensors in the source dataset.

Figure 1 illustrates an example of the above process. Assume that there are two datasets $I$ and $II$, each having 2 and 3 sensors respectively, and their similarity scores have been calculated based on the similarity of their attached objects and deployed locations. Given a current sensor feature $X_I$ in the dataset $I$, we need to simulate a sensor feature $X_{Ii}^{*}$ in the other dataset. First, for each sensor in $II$, we need to identify similar sensors in $I$. Assume that the similarity threshold is 0.5, according to the formula in Definition 3, we identify the similar sensor sets $S_J(j = 1, 2, 3)$ for each sensor in the dataset $II$, as $\{s_{1,1}\}$, $\{s_{1,1}\}$, and $\{s_{1,2}\}$. Then the probability on each sensor is the averaged contribution from their similar sensors. The location and object ontologies as well as the sensor similarity calculation can refer to the work [31].

![Figure 1: An example of sensor feature space remapping](image)

With feature remapping, for each dataset, we will not only train a classifier on its own train data but also on the data converted from the training data in the other datasets. This gives rise to Algorithm 1.

**Algorithm 1 SD: Share training Data**

**Require:** a collection of datasets $\{D_1, D_2, ..., D_n\}$, where each dataset $D_i$ is composed of a set of labelled examples $L_i$ and a set of un-labelled examples $U_i$.

1. initialise a classifier $C_i$ for each dataset
2. for $t = 1$ to $T$
3. $L_i \rightarrow L_i \cup \{L_j' | L_j' = \text{remap}(L_j,i), j \neq i, j \in [1,n]\}$
4. use labelled examples in each dataset $L_i'$ to train its classifier $C_i$ and estimate an uncertainty measure $\mu_i$
5. randomly create a set of $\{U_1', U_2', ..., U_n'\}$ from un-labelled examples $U_i$
6. use $C_i$ to label $U_i'$ and select $k$ most confident and certain examples $K_i$
7. $L_i \rightarrow L_i \cup K_i$
8. $U_i \rightarrow U_i - K_i$
9. end for

**Algorithm 2 SC: Share Classifiers**

**Require:** a collection of datasets $\{D_1, D_2, ..., D_n\}$, where each dataset $D_i$ is composed of a set of labelled examples $L_i$ and a set of un-labelled examples $U_i$.

1. initialise a classifier $C_i$ for each dataset
2. for $t = 1$ to $T$
3. use labelled examples in each dataset $L_i$ to train its classifier $C_i$ and estimate an uncertainty measure $\mu_i$
4. randomly create a set of $\{U_1', U_2', ..., U_n'\}$ from un-labelled examples $U_i$
5. for each unlabelled example $u \in U_i'$
6. $\text{prob} \leftarrow C_i$ classifies $u$ and returns its class probabilities
7. if $\mu_i$ evaluates $\text{prob}$ to be uncertain then
8. use $\{C_1, ..., C_n\}$ to classify $u$ and integrate the results
9. end if
10. end for
11. select $k$ most confident and certain examples $K_i$ from $U_i'$
12. $L_i \rightarrow L_i \cup K_i$
13. $U_i \rightarrow U_i - K_i$
14. end for

to label, and (2) how to integrate results from multiple classifiers. To address the first question, we will employ the most common uncertainty sampling strategies from active learning, which are least confidence, margin sampling, and entropy, as described in Equation 1, 2, and 3 respectively [20].

1. **least confidence:**
   \[
   \phi(x)_{LC} = \arg\max_x 1 - P_\theta(\hat{y}|x) \tag{1}
   \]
   where $\hat{y} = \arg\max_x P_\theta(y|x)$ denotes the class label assigned with highest posterior probability. A larger value of $\phi(x)_{LC}$ denotes a higher uncertainty.

2. **margin sampling:**
   \[
   \phi(x)_M = \arg\min_x P_\theta(\hat{y}_1|x) - P_\theta(\hat{y}_2|x) \tag{2}
   \]
   which computes the difference between two highest posterior probabilities. A smaller value of $\phi(x)_M$ (smaller margin) denotes a higher uncertainty, suggesting it is hard to differentiate two most probable class labels.

3. **entropy:**
   \[
   \phi(x)_E = \arg\max_x - \sum_i P_\theta(\hat{y}_i|x) \log P_\theta(\hat{y}_i|x) \tag{3}
   \]
where entropy of each instance measures uncertainty of the corresponding set of posterior probabilities for all class labels. A larger value of \( \phi(x)_E \) denotes a higher uncertainty.

With the uncertainty sampling strategies, we can determine which example is not certain to be labelled from its current classifier. Then we will perform a feature space remapping that converts it to an example in the other datasets to allow the other classifiers to label. Once the other classifiers complete the inference, we will need to integrate their results to make a final decision. There are existing approaches to combine classifiers; for example, ensemble methods like boosting, bagging, and stacking are classic ways to do. However, we are constrained by the size of training data, so we propose a uncertainty-driven approach. It selects the most certain activity label from each classifier in the other datasets and the process is described in Algorithm 3. To do so, we will reuse the above uncertainty sampling strategies on the inference results and calculate their uncertainty score with the above formulas.

Algorithm 3 CC: Combine Classifiers

```
Require: an unlabelled example \( u \)
Require: Classifiers \( \{C_1, C_2, ..., C_n\} \) and uncertainty measure \( \{\mu_1, \mu_2, ..., \mu_n\} \) for each dataset \( i \in [1, n] \)
1: initialise \( res \) to store the inference results
2: for each classifier \( C_i \) do
3: \hspace{1em} \( prob \leftarrow C_i \), classifies \( u \) and returns its class probabilities
4: \hspace{1em} if \( \mu_i \) evaluates \( prob \) to be certain then
5: \hspace{2em} add \([cls, conf, us]\) to \( res \), where \( cls \) is the inferred class label, \( conf \) is the probability on \( cls \), and \( us \) is the uncertainty score.
6: \hspace{1em} end if
7: end for
8: if \( res \) is not empty then
9: \hspace{1em} return \([cls, conf]\) with the lowest uncertainty score from \( res \)
10: end if
```

### V. Evaluation Methodology

We hypothesise that SD and SC Algorithms can significantly improve activity recognition accuracies when the available training data is very little. To validate this hypothesis, we design the following methodologies.

#### Datasets

We use three real-world smart home datasets that capture typical activities and more importantly they represent common types of smart home datasets in terms of the number of sensors, different spatial layouts, and the degree of inherent noise. We believe that experimenting on these datasets gives us a comprehensive view of the effectiveness of the proposed technique. The three datasets were collected by the University of Amsterdam (named House A, B, and C respectively in the following) from three real-world, single-resident houses which were instrumented with wireless sensor networks [9]. These three datasets recorded the same set of 7 activities, including leaving the house, preparing breakfast or dinner, and sleeping. These three houses are deployed with only binary sensors, whose reading indicates whether or not a sensor fires. More specifically, the House A dataset consists of 14 state-change sensors attached to attached to household objects like doors, cupboards, and toilet flushes, while the other two datasets contain more than 20 sensors, including reed switches to measure whether doors and cupboards are open or closed; pressure mats to measure sitting on a couch or lying in bed; mercury contacts to detect the movement of objects (e.g., drawers); passive infrared to detect motion in a specific area; float sensors to measure the flush of toilet.

#### Accuracy

We consider two metrics for accuracy: the overall accuracy – the ratio of the learnt labels being the same as the true labels, and the class accuracy – the averaged ratio of the learnt labels being the same as the true labels in each class.

\[
A_o = \frac{|\text{pred}_\text{label} == \text{true}_\text{label}|}{N}
\]

\[
A_c = \frac{\sum_{c=1}^{C} (|\text{pred}_\text{label}_c == \text{true}_\text{label}_c|)}{\sum_{c=1}^{C} (N_c)}
\]

where \( C \) is the number of classes and \( N_c \) is the number of test examples in each class \( c \).

#### Process

We evaluate the algorithms across different ratios of training data, from 0.1% to 80%. 0.1% is the extreme training data percentage for the chosen datasets; that is, the training data for each dataset can only contain 1 or 2 examples. To validate our hypothesis, we want to observe the effectiveness of the proposed SD and SC Algorithms when training data is very little and find to what percentage of training data both algorithms need to be able to produce satisfactory accuracies.

For each given ratio \( r \) of training data, we randomly generate \( r \times N_i \) number of instances in the \( i \)th dataset to be annotated, run an algorithm on all the datasets, and calculate the averaged class accuracies. The algorithms include the proposed SD and SC Algorithms as well as the baseline approach; that is, train a classifier with training data of each dataset and test on its test data, without any sharing data or classifiers. We run 100 iterations for each ratio, test all these algorithms on each iteration, and average their accuracies.

#### Classifiers

We employ a collection of the state-of-the-art classification algorithms: Naive Bayes (NB), Random Forest (RF), and Support Vector Machine. None technique consistently significantly outperforms the others. Also a classifier is not the focus of our approach as both SD and SC Algorithms work on top of any state-of-the-art classifiers. In the next section, we will report the results on NB and RF, for the following reasons. These two algorithms represent two different styles of classifiers: NB is a simpler model and consumes less training data to produce good classification accuracies, while RF has more capacity in building complex models but requires much more training data. We want to see how these two algorithms work when the training data is little and how SD and SC Algorithms can enhance their performance in this situation.
Uncertainty sampling strategies

We have experimented with the three uncertainty sampling strategies in Section IV-B. For each strategy, we need to estimate the threshold to determine whether inferred class probabilities are uncertain or not. A general way to do is to split the training data into training and validation sets so that we can experiment with different threshold values and test on the validation set, and we choose the best threshold that leads to the highest accuracies. However, we are dealing with very small training data, we cannot afford the validation set. What we have done is that after training a classifier, we calculate the uncertainty scores of each training example and use the mean score as the threshold. For the extreme case with only one training data, we set the default threshold score on the most uncertain situation; that is, evenly distributed class probabilities. Also we allow the uncertainty threshold to be updated during the iterative training phase, so the default threshold only serves at the cold start stage.

In terms of combining classifiers, we have evaluated CC Algorithm and compared with the classic ensemble methods such as majority voting (MV) and highest confidence (HC) and Dempster-Shafer Theory (DST) to integrate imperfect evidences from distinct sources [14]. The MV, HC, and DST have similar results, all worse than SC Algorithm as they are more likely to be affected by the weaker classifiers. CC Algorithm with the information entropy uncertainty sampling strategy produces the highest performance.

Top $k$ example selection

In both SD and SC Algorithms we will need to choose top $k$ examples for the next training iteration. We have experimented with different numbers 2, 5, and 10. Both top 2 and 5 produce better accuracies as they only focus on best examples while filtering less confident ones. For the sake of performance, we will choose top 5, which leads to faster convergence.

VI. RESULTS AND DISCUSSION

This section will discuss the evaluation results and validate our hypothesis. Figure 2 presents the comparison of overall and class accuracies between Baseline, SD, and SC Algorithms on each dataset.

SD and SC Algorithms improve both overall and class accuracies

When the ratio of training data is small; i.e., from 0.1% to 10%, and perform comparably with the baseline approaches afterwards. To quantify the improvement, we perform significance tests on the performance of SD, and SC Algorithms over the Baseline Algorithm with their corresponding base classifiers respectively. For each ratio of training data, we run Welch’s t test on the accuracies of 100 iterations and calculate the p-values, which are presented in Figure 3. With a null hypothesis H0 of SD, and SC providing no improvement over B in recognising activities, and an alternative hypothesis H1 of SD, and SC displaying improvement, we select a standard significance level of 90% for the test, meaning that if the p-value in the test result is smaller than 0.1, we reject the null hypothesis and accept that there is a statistically significant improvement. The result in Figure 3a shows that when the training ratio is less than 1%, SD has significantly improved the overall and class accuracies (not on RF) with the p-values all close to 0. The main reason is that accumulating training data from multiple sources does provide better coverage. Again in the extreme situation, if training data from each dataset contains one example from different classes, then collecting them together will cover three classes. When the training data is small, SD Algorithm with NB works much better than RF; that is, the averaged class accuracies of 55% with NB and of 17% with RF on House A when the ratio of training data is 0.1% in Figure 2. The reason is that RF often needs more training data to compose a number of decision tree classifiers and less sensitive to small change in training data, while NB is built on a much simpler model and can adapt well to small change in training data.

SD Algorithm is less sensitive to the amount of training data; for example on House A, the averaged class accuracy is 55% at the lowest ratio 0.1%, and 59% at the highest ratio 80% in Figure 2. When the ratio of training data increases, SD Algorithm performs worse than B and SC Algorithms. It suggests that when training data is sufficient, sharing data across different datasets does not help, but adds noise to the classifiers in each dataset. Partly it is because each user tends to have their own way or routine of performing certain activities; for example, interacting with different objects. Sharing data can enhance activity coverage, but not necessarily improve identifying activities for individual users.

SC Algorithm needs more training data than SD Algorithm to perform well, as there is no significant improvement when the training ratio is very small; i.e., less than 0.4% in Figure 3b. The significant improvement on SC (with NB) is from 0.4% to 20%. First of all, SC Algorithm still trains its own classifier on training data from its own dataset; that is, the training data size is not enlarged like SD Algorithm. But when the training ratio increases, SC has better improve-
Fig. 2: Comparison of averaged overall and class accuracies between baseline, SD, and SC Algorithms, showing SD and SC Algorithms can improve recognition accuracies when training data are small.

Fig. 3: Significance test on the performance between baseline, SD, and SC Algorithms, showing SD and SC Algorithms can improve recognition accuracies when training data are small and SD works better than SC Algorithm.

ment than SD. Secondly, the small training data significantly undermines the estimation of the uncertainty threshold. This becomes a more severe problem on RF than NB, as both classifiers produce very different class probabilities in the face of uncertainty. For example, under the extreme situation that there is only one example in the training data, saying it belongs to the 2nd class, for any test example, RF will derive the class probabilities \([0, 1, 0, 0, 0, 0]\); that is, no matter which class the test example might belong, it will assign the highest probability 1.0 to the observed class in the training. On the contrary, NB will derive evenly distributed class probabilities \([0.14, 0.14, 0.14, 0.14, 0.14, 0.14]\). As introduced before, we hardcode the uncertainty threshold as the most uncertain class probabilities, which can detect an uncertain example for labelling on NB but not on RF. This raises a question on what heuristics we should use to better estimate uncertainty measures, which will be one of our future work.

Fig. 4: Comparison of the recognition accuracies of different ways of sharing classifiers on House A
Due to this problem, we are also wondering what if we always share classifiers; that is, for each unlabelled example, we query all the classifiers and integrate their results. The results are presented in Figure 4, where SC2 refers to the always-sharing algorithm. With always sharing classifiers, SC2 Algorithm improves on RF, but not on NB. The results confirm our analysis that the poorer performance on RF is mainly to do with the uncertainty evaluation.

Comparison with active learning Here we will also compare the performance of SD and SC Algorithms with active learning in Section III-B, one of the most applied semi-supervise learning techniques in activity recognition. The performance of active learning with the information entropy uncertainty sampling strategy is presented in Figure 5, including the times of queries to acquire labels on uncertain examples from human operators and the overall and class accuracies. Compared to the results in Figure 2, active learning achieves better accuracies, however, it requires query 10% or 20% of test data to human operators, which makes the actual amount of training data is actually much higher than the specified. Also active learning assumes the always availability of true labels for each user’s own sensor data. On the one hand, this allows annotating sensor data for individual users and thus improves their own activity recognition model. On the other hand, it has better coverage of the activity space than our proposed methods in that the labels that we share in SD and SC are constrained by their availability in the other datasets’ training data. For example, if all the datasets only contain “having a meal” and “sleeping” activities altogether, then our techniques will not be able to detect any activities other than these two, but the active learning technique will still be able to learn the others such as “taking shower” or “having drink” because of the always availability assumption.

VII. Conclusion and Future Work

This paper aims to address the problem on the scarcity of activity annotations by leveraging annotations across different datasets that can have different sensing technologies, different sensor deployment, and different users, as long as they share the same mission with compatible sensor features and activities of interest. The goal of this paper is to explore the solution space to address the problem of SLearn and validate whether the proposed approach can improve activity recognition accuracies given a small ratio of training data and if so, to what degree. To do so, we have proposed two approaches: SD Algorithm – accumulating a larger training data pool by semantically remapping training data from each other, and SC Algorithm – acquiring and integrating activity labels from classifiers on the other datasets. We have presented the general skeletons of these two types of algorithms, and plugged in specific classifiers, feature space remapping techniques, and ensemble methods.

The evaluation results have consistently demonstrated a significant improvement on activity recognition accuracies when training data is small; for example, each dataset only contributes 2 or 3 annotated examples (i.e., the ratio of training data is 0.4%), the averaged class accuracies can reach around 66%, 52%, and 55% on House A, B, and C respectively. These algorithms can save a smart home system a large amount of time and effort on collecting ground truth. They can also potentially support lifelong learning of activity models as long as users from different smart home environments can contribute a few examples of their new activities, which can be either shared by SD Algorithm with the system in the other smart home environments or integrated by SC Algorithm when similar sensor data are reported in the other home environments.

To note that SD and SC Algorithms are not committed to any of the experimented techniques, and can be configured with any other suitable ones. In the future, we will look into other more sophisticated ensemble methods such as Bayesian classifier combination [10], which have demonstrated strength in combining imperfect classifiers [21]. Also we will look into how to use other heuristics to better estimate uncertainty thresholds when the training data is limited, which has significantly affected the performance of certain classifiers like RF in our experiment. To further test the generality of the proposed techniques, we will attempt to working on the other types of application domains such as activities in the office building and outdoor activities, and the other types of data such as accelerometer or physiological data.
REFERENCES


