**XLearn: Learning Activity Labels Across Heterogeneous Datasets**

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Sensor-driven systems often need to map sensed data into meaningfully-labelled activities in order to classify
the phenomena being observed. A motivating and challenging example comes from human activity recognition
in which smart home and other datasets are used to classify human activities to support applications such as
ambient assisted living, health monitoring, and behavioural intervention. Building a robust and meaningful
classifier needs annotated ground truth, labelled with what activities are actually being observed – and
acquiring high-quality, detailed, continuous annotations remains a challenging, time-consuming, and error-
prone task, despite considerable attention in the literature. In this paper we use knowledge-driven ensemble
learning to develop a technique that can combine classifiers built from individually-labelled datasets, even
when the labels are sparse and heterogeneous. The technique both relieves individual users of the burden of
annotation, and allows activities to be learned individually and then transferred to a general classifier. We
evaluate our approach using four third-party, real-world smart home datasets and show that it enhances
activity recognition accuracies even when given only a very small amount of training data.

CCS Concepts: • Computing methodologies → Ensemble methods; • Human-centered computing → Ambient intelligence; Ubiquitous and mobile computing systems and tools;

Additional Key Words and Phrases: Human activity recognition, ensemble learning, transfer learning, clustering,
smart home

ACM Reference Format:

1 INTRODUCTION

Over the last decades more and more smart home and intelligent environment prototypes and
real-world testbeds are emerging, which have demonstrated promising values in the application
domains of health [33], energy [9], and workforce. For example, care homes are deploying sensorised
assisted living platforms to identify the elderly’s daily routine in order to provide personalised
care services, and smart home technology industries aim towards energy-efficient solutions for
air purification or heating configuration based on detected human behaviours such as cooking
and sleeping. Sensor-based human activity recognition is the key enabling technology for these
applications, integrating and abstracting low-level sensor data into high-level activities to which

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activity-aware applications can adapt their behaviours. The ability to correctly identify and predict users’ activities underpins the success of these applications.

Significant progress has been made in activity recognition over the past years with the support of a large number of modern data-driven techniques [2, 38], including Hidden Markov Models, Conditional Random Fields, Support Vector Machine, and more recently deep neural networks [27]. To build a robust activity recognition model, most of these existing techniques require a large volume of training data; that is, sensor data annotated with activity labels. However, the key challenge facing the current activity recognition community is the lack of sufficient training data. On the one hand, it often requires a lot of time and effort to annotate sensor data. It either relies on users’ constant self-report on what they are doing, or records users’ activities via videos which are later annotated by the others. On the other hand, there is never enough training data, as users might change their routine over time; that is, they may perform new types of activities such as playing Kinnect fitness games to be more active – and in doing so change both the activities they do and the ways in which they do them. Or users’ health conditions degrade due to ageing or onset of a certain disease, which can change their daily routine. Therefore annotation approaches that require highly intensive effort and commitment are neither scalable nor sustainable for a large number of users over a long period of time.

In this paper, we propose a novel approach, called XLearn, to reduce the annotation burden on individual users via cross learning on scarcely- and partially-annotated data from multiple users, to achieve satisfactory activity recognition accuracies. The hypothesis is that, as long as each user contributes a very small number of labelled examples (even though these examples might not cover a complete set of activity types), a cross learning approach will learn annotated examples across all the users and be able to build an activity recognition model for each user to recognise all activities. This is illustrated in Figure 1.

**Fig. 1.** An example of XLearn scenario. We have two users living in two residential settings deployed with different sensors. Each user has been annotated with different activities. XLearn aims to combine their sensor data and activity annotations to learn all these activities on all the users.

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in different home settings deployed with a different number of sensors in different types of sensing technologies) and may have different but semantically related output labels (i.e., source and target domains might target different types of activities, or use different labels for the same activities).

**XLearn** is a knowledge-driven ensemble learning technique, built on generic smart home ontologies to enable computationally efficient feature and activity space remapping. In our previous work, **SLearn** [37], we have achieved preliminary, promising results on using the ontologies to remap feature spaces from heterogeneous datasets, but the performance of **SLearn** needs significant improvement. In this paper, to enable robust integration of inferred activity labels from a diverse collection of datasets, we employ a stacking ensemble and incrementally update the ensemble classifier and its base classifiers over time to improve the accuracies of activity recognition. Also **SLearn** assumes that all the datasets share the activity labels, which can be possible if the same organisation deploys an activity monitoring system in many different houses and is more likely to monitor a similar set of activities. However, **XLearn** works on a more general assumption where each dataset can have a different set of activities as long as these activities are semantically related.

We have evaluated **XLearn** on four real-world smart home datasets, which exhibit high heterogeneity in terms of environmental layouts, sensing technologies, and activities. The results demonstrate that **XLearn** has achieved satisfactory recognition accuracies with a small number of training data (i.e., 2 from each dataset), better than the classic activity recognition approach – only using each user’s own data for training. While we have developed and exercised our technique on smart-home data, the underlying approach promises to be applicable to any sensor-driven activity-recognition problem.

The rest of the paper is organised as follows. Section 2 scopes the problem that **XLearn** aims to address with formal definitions and an illustrative example. Section 3 reviews the mainstream approach on addressing the scarcity of labelled data and identifies the difference between these other techniques and **XLearn**. Section 4 describes the **XLearn** approach, and Section 5 introduces the evaluation methodology and experiment setup. Section 6 performs an evaluation and discusses the technique’s strengths and limitations, and Section 7 concludes the paper.

## 2 PROBLEM STATEMENT

In this section, we will identify the research problem that the **XLearn** tries to address, through a concrete illustrative example and a formal definition. Take Figure 1 as an example scenario, which is activity recognition in a smart home environment deployed with binary event-driven sensors. Assume there are two home settings: House A and W, each with different spatial layouts and hosting different users. Each environment is instrumented with a different number of binary event-driven sensors of different sensing technologies, including infra-red passive motion detectors [8, 16], door or window switch sensors, object sensors, and pressure sensors [16, 22]. These sensors collect data about a resident’s presence in the environment, and their interactions with everyday objects, which will help to interpret what the resident is doing [39]. Examples of sensor events and labelled activities are presented in Figure 1.

Based on the syntax and terminology from transfer learning [26], we first give preliminary definitions of problems that the **XLearn** approach aims to address.

**Definition 1.** Let a dataset \( \mathcal{D} \) consist of a domain \( D \) and a task \( T \), where

1. a domain \( D \) is a two-tuple \((\chi, P(\chi))\). \( \chi \) is the \( m \)-dimensional feature space of \( D \) and \( P(\bar{x}) \) is the marginal distribution where \( \bar{x} = [x_1, ..., x_m] \in \chi \).
2. a task \( T \) is a two-tuple \((\mathcal{Y}, f())\) for a domain \( D \), \( \mathcal{Y} \) is the label space of \( D \), which is a collection of possible class labels \( \mathcal{Y} = \{y_1, y_2, ..., y_{N_c}\} \). \( f() \) is an objective predictive function for \( D \).
(3) within this dataset, each pair \((\bar{x}, y) \in \chi \times \mathcal{Y} \cup \{\Theta\}\), where \(\Theta\) represents unknown or unlabelled.

That is, some instances in the feature space have labels; i.e., \((\bar{x}, y) \in \chi \times \mathcal{Y}\) and \(f(\bar{x}) \in \mathcal{Y}\), while the other instances do not; i.e., \((\bar{x}, \Theta) \in \chi \times \{\Theta\}\) and \(f(\bar{x}) = \Theta\).

Take an example of the datasets in Figure 1, an instance \(\bar{x}\) is represented as a feature vector \([x_1, x_2, ..., x_n]\) in \(\chi\), where \(n\) is the number of sensors being deployed and \(x_i\) indicates the ratio of the \(i\)th sensor reporting an event during a certain time interval (e.g., every 30 or 60 seconds); that is, \(x_i = N_i / N\), where \(N_i\) is the number of events reported by the \(i\)th sensor and \(N\) is the total number of events being reported in an interval. This feature representation is similar to the numerical format to represent binary sensor data in a vector [4], and the only difference is that we normalise the number of activation with the total sensor events in each interval. Another way to represent sensor feature vector values as 0 or 1 to indicate whether a sensor is activated or not. This binary representation can be difficult to distinguish activities that activate the same set of sensors; for example, when a user is preparing for breakfast and dinner, he/she might activate most of the sensors in the kitchen.

The label space \(\mathcal{Y}\), for example in House A, consists of 'prepare breakfast' and 'get drink'. An unknown label on sensor data is marked as \(\Theta\). The predictive function \(f\) here is to classify sensor features \(\bar{x}\) into an activity label in \(\mathcal{Y}\).

**Definition 2.** Let \(\mathcal{DS}\) be a collection of datasets \(\{D_1, D_2, ..., D_{N_D}\}\), where each dataset has different domains (i.e., \(\forall i, j \leq N_D, i \neq j, \chi_i \neq \chi_j\) and \(P_i \neq P_j\)) and has different tasks (i.e., \(\mathcal{Y}_i \neq \mathcal{Y}_j\) and \(f_i \neq f_j\)). \(XLearn\) aims to assign a label to each unlabelled instance in the dataset; that is, \(\forall \bar{x} \in \chi_i\), there exists a label \(y \in \mathcal{Y}_i \cup \mathcal{Y}_2 \cup ... \cup \mathcal{Y}_{N_D}\). To make this work we make two assumptions:

1. **Feature space remapping** – the feature spaces between the datasets are comparable and can be remapped between each other. That is, there exists a mapping function \(\theta_{i \rightarrow j} : \chi_i \rightarrow \chi_j\) that remaps each instance \(\bar{x}\) in \(\chi_i\) to an instance \(\bar{x}'\) in \(\chi_j\).
2. **Label space remapping** – similarly, there exists a mapping function \(\theta_{i \rightarrow j} : \mathcal{Y}_i \rightarrow \mathcal{Y}_j\) that remaps each label \(y\) in \(\mathcal{Y}_i\) to a label \(y'\) in \(\mathcal{Y}_j\), and \(y\) and \(y'\) can be in different terms but are semantically related.

As we can see in Figure 1, each dataset features different spatial layouts and sensor deployments, so they have different feature spaces in terms of feature dimension and semantics and as well as their marginal distributions; i.e., the frequency of a sensor being activated.

In addition, each dataset targets different collections of activities; that is, different numbers of activities, and different labels for the same or semantically similar activity such as 'meal preparation' in House W and 'prepare breakfast' in House A. It is clear that these two datasets have different domains and tasks with respect to Definition 1. The sensor events in each house are partially labelled; that is, some sensor events have not been annotated with an activity label but with the unknown symbol \(\Theta\) in Figure 1. The task of \(XLearn\) is to complement each dataset’s annotation and recognise all these 4 activities on unlabelled sensor data in both datasets.

We present \(XLearn\) as a promising technique to improve activity recognition on scarcely labelled datasets, all of which can have different feature spaces and label spaces, and do not have to be fully labelled. Thus, this technique is not subject to heterogeneous sensor deployments and activity sets in different environments, which addresses the key challenge in transfer learning of human activity recognition in the real-world applications. To make the learning effective, \(XLearn\) builds on an assumption that there exists simple ontologies that semantically relate the feature spaces across different datasets and as well as their label spaces.
3 RELATED WORK

Activity recognition has been an active research topic in the last decade, and a variety of techniques have been proposed [38]. Among them, different approaches have been designed to address the scarcity challenge of activity annotation, including knowledge-driven techniques, unsupervised learning, semi-supervised learning, and transfer learning. In the following, we will compare and contrast these approaches with XLearn.

3.1 Knowledge-driven Techniques

Ontologies are one of the main knowledge-driven approach in activity recognition [38]. Chen et al. [5] present a comprehensive smart home ontology for daily activity recognition. For example, a ‘making tea’ activity can be described with a collection of property-value pairs, including hasContainer, hasTeabag, and hasFlavour. In general, the knowledge-driven approaches often require heavy knowledge engineering effort; for example, specifying how each activity is related to objects or events that can be sensed from sensors. Such activity specification can be error-prone or difficult to accommodate different variations or patterns of an activity.

Research has been carried out to integrate knowledge with machine learning techniques. Riboni et al. [30] propose a hybrid approach of ontological and probabilistic reasoning for activity recognition. It combines ontologies with Markov Logic Network to reason temporal relationships between sensor events and activities; for example, ‘turning on the oven’ cannot belong to an activity instance ‘meal preparation’ if their temporal distance is too far apart; e.g., over two hours. Ye et al. [39] have used a lightweight ontology to specify necessary conditions of an activity. Based on this ontology, a collection of machine learning techniques including sequential mining and clustering are applied to enable unsupervised learning on activity recognition.

XLearn applies ontologies to support feature space remapping, but itself is better than an knowledge-driven activity recognition approach, it does not need a detailed specification on activities, which often requires a lot of knowledge engineering effort and can be error-prone. The problem that XLearn aims to address is different from a classic activity recognition problem; that is, XLearn tries to shared partially learned activity models from multiple datasets to complement the learning due to insufficient training data, rather than learning activity labels on one dataset.

3.2 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 [14]. Rashidi et al. propose a method to discover the activity patterns and then manually group them into activity definitions [28]. 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, in our previous work, we have combined the sequential mining and clustering algorithms to discover representative sensor events for activities [39]. Different from the work in [28], they have applied the generic ontologies to automatically map the discovered sensor sequential patterns to activity labels through a semantic matching process [39]. Yordanova et al. have also applied domain knowledge in rule-based systems to generate probabilistic models for activity recognition [19, 42].

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 [25, 36, 41, 43]. During the reasoning process, the mined objects are mapped to sensor events and an appropriate activity will be recognised.

XLearn targets a problem more challenging than a classic activity recognition problem where for each dataset we train a model with labelled instances and recognise unlabelled instances. In XLearn, the training data is assumed to be incomplete in that it might not cover all the activities of interest. XLearn is not an unsupervised learning technique as it still relies on labelled training data, but the labels can come from different datasets. In this way, XLearn shares the annotation cost across a large number of users, which can significantly reduce the cost on individual users.

3.3 Semi-supervised Learning

Active learning, also called ‘query learning’, is a subfield of machine learning motivated by the scenario where there is a large amount of unlabelled data and a limited but 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 [31].

Stikic et al. have attempted active learning and cotraining to recognise daily activities from accelerometers and infra-red sensors [32]. 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 classes [1]. The annotated instances are then used to iteratively update a HMM to infer daily activities in a home setting. 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 [6]. Similarly, Hossain et al. combine the uncertainty measure and Silhouette coefficient to select the most informative instances as a way to discover new activities [15].

XLearn is better than active learning [31] in that it will not query a user or a human operator, but learns labels from the other datasets, which again reduces the annotation burden on each user. Another commonly applied semi-supervised learning technique is cotraining, proposed by Blum and Mitchell in 1998 has been one classic approach able to boost performance of a learning algorithm by leveraging a large number of unlabelled examples [3]. 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 cotraining algorithm and its variations have been recently applied in the multi-view human activity recognition in smart home environments [13]. 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.

Cotraining works on multiple views from the same data while the XLearn approaches need to work on multiple datasets that share the sensing mission; that is, compatible sensor features and activities. Here XLearn novelty applies the cotraining principle in transfer learning.

3.4 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) [26]. Maekawa et al. [23] 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.
Zheng et al. [43] 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. 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.

van Kasteren et al. [34] 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. [29] 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. [12] 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.

XLearn is most relevant to but not the same as transfer learning [26] as it targets a research problem that challenges the assumptions of most transfer learning techniques. Here 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 may 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 XLearn. 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 with the advantage of reducing the impact of sensing technology, deployment, and individual activity routine on the effectiveness of transfer learning.

3.5 Comparison and Summary

In summary, Table 1 has contrasted and compared XLearn with the existing activity recognition techniques. XLearn has worked on a different assumption: instead of learning an activity model from one dataset, it cross learns the activity models between different datasets by leveraging an activity model from each dataset and complementing each other’s insufficiently annotated training data. It does not require all the activity labels are annotated in each dataset. To do so, we design algorithms to bridge and learn the difference between their feature and label spaces.

4 PROPOSED APPROACH

We hypothesise that when each dataset contributes a small number of labelled examples (even though these examples might not cover a complete set of activity types), XLearn will be able to cross learn the annotated examples and improve activity recognition accuracies. XLearn is a knowledge-driven ensemble learning technique, whose main design points are illustrated as follows.

(1) The key enabler for sharing these datasets is the feature space remapping and label space remapping. We will propose a generic knowledge-driven approach in Section 4.1 and justify the advantage of this approach by comparing it against a data-driven approach.

(2) As we aim to reduce the annotation burden, we will reduce the amount of the training data as much as possible. However, a small amount of training data might prevent from building...
Table 1. Comparison between XLearn and state-of-the-art activity recognition techniques

<table>
<thead>
<tr>
<th>Techniques</th>
<th>Datasets</th>
<th>Labelled</th>
<th>Learning</th>
</tr>
</thead>
<tbody>
<tr>
<td>Supervised Learning</td>
<td>Single dataset</td>
<td>Fully</td>
<td>Train on well-annotated dataset and recognise newly incoming sensor data [38]</td>
</tr>
<tr>
<td>Knowledge-driven</td>
<td>Single dataset</td>
<td>Not</td>
<td>Specify each activity on the knowledge model and infer activity labels from sensor data [5]</td>
</tr>
<tr>
<td>Hybrid model</td>
<td>Single dataset</td>
<td>Fully</td>
<td>Train the complex correlation between sensor data and activities, leverage the knowledge model in guiding and constraining the training process with activities specified on the knowledge model [30, 39], and infer activity labels from sensor data</td>
</tr>
<tr>
<td>Unsupervised learning</td>
<td>Single dataset</td>
<td>Not</td>
<td>Learn the latent structure of sensor data and assign activity labels to the structures accordingly based on the knowledge specified or learnt [19, 28, 39, 42]</td>
</tr>
<tr>
<td>Semi-supervised learning</td>
<td>Single dataset</td>
<td>Partially</td>
<td>Train on a small amount of training data, incrementally update the model with self-learnt labels [13, 32] or acquired labels through active learning [1, 6, 15, 32]</td>
</tr>
<tr>
<td>Transfer learning and domain adaptation</td>
<td>2 datasets with homogeneous or heterogeneous feature spaces</td>
<td>Fully</td>
<td>Learn an activity model on one dataset (called the source) and adapt or transfer the model to another dataset (called the target), with a general assumption that the source dataset is annotated with all the activity labels [12, 23, 29, 34, 43]</td>
</tr>
<tr>
<td>SLearn</td>
<td>Multiple (≥ 2) datasets with heterogeneous feature spaces</td>
<td>Partially</td>
<td>Shared learning activity labels from datasets via either sharing the training data or the classifier on each dataset [37]</td>
</tr>
<tr>
<td>XLearn</td>
<td>Multiple (≥ 2) datasets with heterogeneous feature and activity spaces</td>
<td>Partially</td>
<td>Cross learning activity labels from datasets in an ensemble classifier that takes into account the classification performance and output of each dataset’s activity model, and latent structures of each dataset</td>
</tr>
</tbody>
</table>

As each dataset may contain a subset of activities of interest, we need to integrate activity labels inferred from other datasets to complete the activity set. An ensemble classifier is a natural choice for integration and has demonstrated beneficial in analysing complex datasets [18]. Because we deal with very little training data, the classifier on each dataset can be very weak. To make the best of these weak classifiers, we extract a range of meta-features to characterise inference results from each dataset’s classifier and build a stacking ensemble to learn the correlations between these meta-features and the target activity label. This will be described in Section 4.3.

Figure 2 illustrates the above workflow and in the following we will detail each of the processes.

### 4.1 Semantic model

Underlying XLearn is the semantic model of sensor features and activity labels, based on which we can remap the sensor feature space and activity output space from one dataset to another.

#### 4.1.1 Smart home ontologies

There are different feature remapping strategies described in [26]. Feuz et al. [12] 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 environment.
Cluster labeling
Partition data into different subsets and use labelled instances to label unlabelled instances

Ensemble labelling
Acquire the most likely label by integrating inferred labels from all the datasets

Initial Partially-Labelled Datasets

Final Fully-Labelled Datasets

Fig. 2. Workflow of XLearn. At the top are a set of datasets labelled as #1, #2, ..., #ND, each of which contains a set of labelled instances (marked in green, plum, and blue) and a set of unlabelled instances (marked in grey). Cluster labelling will label instances that are semantically close to the labelled instances. This will help expand the existing training set and build a more robust classifier on each dataset. An ensemble is employed to learn the correlation of feature space in each dataset and the combined activity labels from all the datasets. Then the ensemble labelling process integrates the inference results to select the most confident label for each remaining unlabelled instance. This ensemble labelling process will run iteratively and incrementally until all the instances are labelled. The assigned labels can come from different datasets (as shown at the bottom of the figure) 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 adopt a knowledge-driven feature mapping based on smart home ontologies [39] which has demonstrated generality across heterogeneous smart home datasets. The principle of knowledge-driven 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 a hierarchy. Figure 3
presents part of object and location ontologies. For example, Door ⊑ MovableBarrier and Bedroom
⊑ SleepingArea.

Based on the hierarchy, we can measure the similarity between domain concepts [35]. The
approach works by finding the least common subsumer (LCS) of the two input concepts and
computing the path length from LCS up to the root node. LCS is the most specific concept that
both input concepts share as an ancestor.

**Definition 3.** Let \( c_1 \) and \( c_2 \) be two concepts organised in a hierarchy. The conceptual similarity
measure between them is:

\[
sim(c_1, c_2) = \frac{2N_l(N_1)N_2}{N_1+N_2},
\]

where \( N_l(N_2) \) are the paths length between \( c_1 \) and \( c_2 \) and the LCS node of \( c_1 \) and \( c_2 \), and \( N_3 \) is the path length between LCS and the root.

For example, to calculate the similarity between Door and Cup, we locate their LCS as Artifact,
calculate three paths: (1) from Door to Artifact; i.e., 3; (2) from Cup to Artifact; i.e., 4; and (3)
from Artifact to the root PhysicalEntity; i.e., 1, and compute the similarity as \( \frac{2+1}{3+4+2+1} = 0.22 \).

**4.1.2 Feature space remapping** In the following, we define both feature and activity space remap-
ning based on these ontologies.

**Definition 4.** Let \( \vec{x}_I = [x_{I,1}, x_{I,2}, ..., x_{I,m_I}] \) be a \( m_I \)-dimensional feature vector from a dataset \( I \) in
Definition 1. A semantics-based feature mapping function is defined as \( \theta_{I \rightarrow I}(\vec{x}_I) = \vec{x}_{II} \), denoted as
\( [x_{II,1}, x_{II,2}, ..., x_{II,m_{II}}] \), where \( \forall 1 \leq j \leq m_{II} \),

\[
x_{II,j} = \frac{\sum_{i \in S_j} x_i \cdot \text{sim}(s_{I,i}, s_{II,j)}}{|S_j|},
\]

\( S_j \) is a collection of sensors in the feature space \( I \) that are similar to the sensor \( j \) in \( II \).

\[
S_j = \{s_I|1 \leq l \leq m_I, \text{sim}(s_{I,l}, s_{II,j}) > \varepsilon\},
\]

\[
\text{sim}(s_{I,l}, s_{II,j}) = (\text{sim}_L(s_{I,l}, s_{II,j}) + \text{sim}_O(s_{I,l}, s_{II,j}))/2
\]

where \( \varepsilon \) is the threshold to choose similar sensors, \( \text{sim}_L \) and \( \text{sim}_O \) are the similarity measure on
location and object concepts that the \( l \)th sensor in \( I \) and \( j \)th sensor in \( II \), and \( w_L \) and \( w_O \) are the
weights on the location and object similarity; i.e., \( w_L + w_O = 1 \).

Based on the above definition 4, a value $\mathbf{x}_{II,j}$ in a converted instance $\mathbf{x}_{II} \in \chi_{II}$ is the weighted average of the ratio 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 ratio of the $j$th sensor reporting events by looking at the probabilities of all of its similar sensors in the source dataset.

Figure 4 illustrates an example of the above process. Assume that there are two datasets $I$ and $II$, each having 3 and 2 sensors respectively, and their similarity scores have been calculated based on the similarity of their attached objects and deployed locations with both $w_O$ and $w_L$ as 0.5\(^1\). Given a current sensor feature $\mathbf{x}_I$ in the dataset $I$, we need to simulate a sensor feature $\mathbf{x}_{II}$ 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 4, we identify the similar sensor sets $S_j (j = 1, 2)$ for each sensor in the dataset $II$ as $\{s_{I,1}\}$ and $\{s_{I,3}\}$. Then the probability on each sensor is the averaged contribution from their similar sensors.

**Fig. 4. An example of sensor feature space remapping**

### 4.1.3 Activity label space remapping

Within the semantic model, we can also remap activity label space. Even though each dataset can be partially labelled, we assume that the designer and developer of each smart home has pre-defined a collection of activities of interest. Each activity can be described with a collection of location concepts where a user often performs this activity and with a collection of object concepts which this activity often involves [39]. Then the similarity between any two activities is weighted similarity between their corresponding location and object concepts.

**Definition 5.** Given an activity $y_{I,i}$ in the label space $\mathcal{Y}_I$, then a semantics-based label space remapping function is defined as $\delta_{I\rightarrow II}(y_{I,i}) = y_{II,j} = \arg \max sim(y_{I,i}, y_{II,j})$.

Given $y_{I,i}$ maps to a collection of location concepts $L_i$ indicating where the activity can be performed and object concepts $O_i$ indicating what everyday objects might be involved in this activity, and similarly $y_{II,j}$ maps to a collection of location concepts $L_j$ and object concepts $O_j$,

$$sim(y_{I,i}, y_{II,j}) = \sum_{C_i \in \{L,O\}} w_C \times sim_C(C_i, C_j)$$

and

$$sim_C(C_i, C_j) = \frac{\sum_{c_i \in C_i} c_i \times sim_{C}(c_i, c_j)}{|C_i| \times |C_j|}.$$

### 4.2 Cluster labelling

A classic approach of dealing with a small amount of training data is leveraging unlabelled data. That is, for each dataset, we train a classifier on its labelled data and then use it to iteratively 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 expand the labelled data pool and iteratively update the classifier.

However, it is difficult to initialise a robust classifier with too little training data. For example, there are two extracted sensor features, on which the main activated sensors are ‘stove’ and ‘microwave’ respectively. Even though these two features are similar, a classifier trained on the

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1 The location and object weight $w_O$ and $w_L$ can be defined differently in different environments and sensor deployments. Here we consider their contribution to sensor similarity is equal. In our experiments, we have not observed the significant impact of different weight settings on the accuracy of XLearn. The results are reported in the supplementary file: https://drive.google.com/drive/folders/1tQkB0ERk74sTmF5aNkdCrCDidqEC67P.
observed feature (i.e., ‘stove’) will not be able to recognise the new feature (i.e., ‘microwave’). Thus
our first task is to expand the training set on each dataset to label all the unlabelled instances that
are similar to the labelled instances. This will allow us to build a more robust classifier to recognise
as many already-labelled activities as possible.

To enable semantic clustering, we define a semantic cosine based on the soft cosine distance
metric but using the semantics of Definition 4:

$$\text{semantic cosine}(a_i, b_j) = \frac{\sum_{i,j} s_{i,j} a_i b_j}{\sqrt{\sum_{i,j} s_{i,j} a_i^2} \sqrt{\sum_{i,j} s_{i,j} b_j^2}}$$

where a and b are two sensor feature vectors, and $s_{i,j}$ are the similarity between the ith and jth
sensor in Definition 4.

With the semantic cosine as the distance metric, we cluster the whole dataset into different
groups and then perform label expansion in each of the subsets. This process has been illustrated in
Algorithm 1 and 2. For each cluster, we collect the activity labels on its data points. If a cluster has
no label, then we leave it for the ensemble labelling stage. If there exists one label within a cluster,
then we spread the same label to all the unlabelled instances in that cluster under the assumption
that the same activity will normally trigger similar sets of sensors and result in instances whose
distance are close. Otherwise, we split this set further into multiple clusters, until all the clusters
have zero or one label, or indivisibly reach the minimum size of 2.

Algorithm 1: label expansion: expanding labels to semantically similar instances

<table>
<thead>
<tr>
<th>input</th>
<th>A dataset $D$ – a collection of instances, part of which are labelled with activity labels</th>
</tr>
</thead>
<tbody>
<tr>
<td>Result</td>
<td>$LC$ – a collection of clusters whose instances have been labelled</td>
</tr>
<tr>
<td>Result</td>
<td>$UC$ – the remaining unlabelled clusters</td>
</tr>
<tr>
<td>$LC$</td>
<td>$\emptyset$;</td>
</tr>
<tr>
<td>$UC$</td>
<td>$\emptyset$;</td>
</tr>
<tr>
<td>$C$</td>
<td>semantic_cluster($D$);</td>
</tr>
<tr>
<td>foreach $c \in C$ do</td>
<td></td>
</tr>
<tr>
<td></td>
<td>split_and_label($c$, $LC$, $UC$);</td>
</tr>
<tr>
<td>end</td>
<td></td>
</tr>
</tbody>
</table>

4.3 Ensemble Learning

After the above cluster labelling process, we build a base classifier on the expanded training set
on each dataset to recognise the existing activity labels. Ensemble learning here is to infer the
most likely activity label for the remaining unlabelled instances in each dataset by integrating the
inferred labels from all the available datasets. To do so, an intuitive way is an ensemble classifier
that aggregates the inference results from each dataset. There exist different types of ensembles [18]
and here we opt for a stacking ensemble for the following reasons. Even though we have expanded
the training set, a base classifier on each dataset can still be weak and their inference results are
strongly biased towards the activity labels occurring in the training set. For example in the extreme
case where there is only one type of activities being labelled in the training set, a Random Forest
classifier will always infer that activity and the posterior probability on that activity will be 1.0.
Thus the simple ensemble classifiers like majority voting will not work well (or at all). To account
for the weakness of the base classifiers, we will consider the notion of reliability of classifiers,
which has often been used in the fusion process [21].

Modal accuracy is one aspect of the reliability of a classifier, which is often determined by the
overall classification accuracies [20]. However, overall accuracies are indiscriminate on different
misclassification cases [21]. Instead we estimate the reliability using macro F1 scores on the training
set, that is, a balanced precision and recall on each activity class. This score indicates how well the
classifier infers membership of each class, and it will also discount the high posterior probability of
a class that dominates the training set.
\textbf{ALGORITHM 2:} split\_and\_label: splitting a cluster and labelling its instances

\begin{algorithmic}
\Statex \textbf{input} : $c$ - a cluster of instances that might or might not be labelled
\Statex \textbf{input} : $\mathcal{LC}$ - a collection of clusters whose instances have been labelled
\Statex \textbf{input} : $\mathcal{UC}$ - the remaining collection of unlabelled clusters
\Statex // collect unique activity labels for a given cluster $c$
\Statex L \leftarrow \text{collect\_unique\_labels}(c);
\If{$|L| == 0$}
\Statex \quad \mathcal{UC} \leftarrow \mathcal{UC} \cup \{c\}
\ElseIf{$|L| == 1$}
\Statex \quad \text{// label all the instances within the group $c$ with the same label in $L$;}
\Statex \quad \text{label}(c, L);
\Statex \quad \mathcal{LC} \leftarrow \mathcal{LC} \cup \{c\}
\EndIf
\ElseIf{$|L| > 1$ and $|g| > \text{size\_min}$}
\Statex \quad \text{// If there exists more than one label in this cluster, then we need to split the cluster further;}
\Statex \quad C' \leftarrow \text{semantic\_cluster}(c);
\Statex \quad \text{foreach} \; c' \in C'
\Statex \quad \quad \text{// label all the instances within the group $c'$ with the same label in $L$;}
\Statex \quad \quad \text{label}(c, L);
\Statex \quad \quad \mathcal{LC} \leftarrow \mathcal{LC} \cup \{c\}
\Statex \quad \text{end}
\EndIf
\end{algorithmic}

Another aspect of the reliability of a classifier is the 	extit{distance to prototypes}, which measures the distance between a current instance and a prototype; i.e., a representative pattern of a class. The intuition is that the closer is the instance to a prototype, the more likely the instance belongs to the same class as the prototype. Here, we define a prototype as each labelled cluster. For each activity class, we find the cluster that is closest to a given instance $\bar{x}$ and record its distance and size. The size can suggest how reliable the cluster is. The distance is calculated as follows:

\[ \text{dist}(\bar{x}, a) = \min_{1 \leq i \leq n_a} \text{dist}(\bar{x}, c_i), \quad \text{and} \quad \text{dist}(\bar{x}, c_i) = \frac{1}{|c_i|} \sum_{\omega \in c_i} \text{dist}(\bar{x}, \omega), \forall \omega \in c_i \]

where $n_a$ is the number of clusters that are labelled with an activity $a$, and $\bar{x}'$ is any example in the $i$th cluster $c_i$. In the end, for an instance $\bar{x}$ we will generate a meta feature vector $\mathcal{F}$, which is defined as $\mathcal{F} = \{\text{pr}(\bar{x}, c_i) | |\text{dist\_sz}(\bar{x}, c_i)| | \leq \text{size} \},$ where (1) $\text{pr} - \text{posterior probability}$, indicating how likely an instance belongs to each activity from the $j$th dataset’s base classifier; (2) $\text{acc}_j - \text{modal accuracy}$, indicating the accuracy of the $j$th base classifier at inferring activities on the training set; and (3) $\text{dist\_sz}_j - \text{distance and size}$ of the closest cluster in each activity class from the $j$th dataset.

To combine these meta features, we use a stacking ensemble to learn the correlations between these features and an activity label, which can be simply phrased as: $\tilde{y} = \mathcal{F}^T \tilde{\omega}$, where $\tilde{\omega}$ is the learnt weight vector on the generated features and $\tilde{y}$ is the posterior probability on each activity category.

To build a stacking ensemble, we will gather all the labelled instances from each dataset and generate the above features using Algorithm 3. Thus, we collect a new training set $\mathcal{DR} \in \mathbb{R}^{m \times n}$ for the stacking ensemble, where $m = 4 \times \sum_{i=1}^{N_i} N_i$, where $N_i$ is the number of activity classes on the $i$th dataset, and $n = \sum_{i=1}^{N_D} N'_L$, where $N'_L$ is the number of labelled instances in the $i$th dataset. The activity space of the ensemble is the union of the activity labels from all the datasets. The ensemble will aim to learn on the meta features, and balance the posterior probabilities and the modal accuracies of each base classifier and the distance to their closest cluster in each activity category between all the datasets.
ALGORITHM 3: generate_meta_feature: generate features for a stacking ensemble

```plaintext
input : \( x_i \) – an instance from the \( i \)th dataset
input : \( L = \{ LC_j \mid 1 \leq j \leq N_D \} \) – a collection of labelled clusters from each dataset
input : \( H = \{ h_j \mid 1 \leq j \leq N_D \} \) – a collection of base classifier models built on the labelled clusters in each dataset
// generate meta features for an ensemble;
\( F = \) null;
for \( j = 1, \ldots, N_D \) do
    if \( i \neq j \) then
        // remap \( \tilde{x}_i \) from \( i \)th to \( j \)th feature space;
        \( \tilde{x}_j = \theta_{i\rightarrow j}(\tilde{x}_i) \)
    end
    // use \( h_j \) to classify \( \tilde{x}_j \) and get the probability distribution;
    \( pr_j = h_j(\tilde{x}_j) \);
    // get the current model accuracy of \( h_j \);
    \( acc_j = model\_accuracy(h_j) \);
    // for each activity type \( a \), find the cluster that is closest to \( \tilde{x}_j \) and collect its size
    \( (d_j, s_j) = closest\_distance(\tilde{x}_j, LC_j) \);
    \( F = concatenate(F, (pr_j, acc_j, d_j, s_j)) \);
end
return \( F \)
```

ALGORITHM 4: ensemble_builder: build a stacking ensemble

```plaintext
input : \( L = \{ LC_i \mid 1 \leq i \leq N_D \} \) – a collection of labelled clusters from each dataset
// generate features for an ensemble;
\( FS = \) \( \emptyset \);
for \( i = 1, \ldots, N_D \) do
    foreach \((\tilde{x}_i, y) \in unfold(D_i)\) do
        \( FS = FS \cup \{\) generate_meta_feature(\( \tilde{x}_i \)), \( y\)\( \} \);
    end
end
// train an ensemble on the meta feature;
\( h' = train(FS) \)
```

Once the ensemble is built, we will perform online learning and update on the ensemble, which is described in Algorithm 5. For each unlabelled instance \( \tilde{x} \) that is randomly sampled from unlabelled clusters in each dataset, we will generate the meta feature and derive the activity label with the likelihood from the ensemble; that is, \( y_{max} \leftarrow \arg \max_{y \in \mathcal{Y}} pr(a = y|\tilde{x}) \).

We rank the instances according to their likelihood, and select the top \( k \) instances to annotate. Following Definition 5, we will remap the predicted labels for these selected instances to the labels in their own activity space. For example in Figure 1, if an instance from House A is predicted as ‘W.meal preparation’ – a label from House W, this label will be remapped to House A as ‘A.prepare breakfast’. Then we use these instances to update the unlabelled clusters by spreading their labels onto the other examples in the same cluster, move these labelled clusters from \( UC \) to \( LC \), and update each base learner and the ensemble with newly-labelled instances. We iterate the above process until all the instances are labelled.

5 EXPERIMENTS AND EVALUATION METHODOLOGY

The main objective of the evaluation is to assess whether \( XLearn \) can achieve accurate activity recognition with limited training data. More specifically, we want to address the following questions:
ALGORITHM 5: ensemble_update: online learning of a stacking ensemble

input : $\mathcal{L} = \{\mathcal{LC}_i | 1 \leq i \leq N_D\}$ - a collection of labelled clusters from each dataset

input : $\mathcal{U} = \{\mathcal{UC}_i | 1 \leq i \leq N_D\}$ - a collection of labelled clusters from each dataset

input : $\mathcal{H} = \{h_i | 1 \leq i \leq N_D\}$ - a collection of base classifier models built on the labelled clusters in each dataset

input : $h'$ - a trained ensemble classifier

while $\exists \mathcal{UC}_i$ NOT empty do
  $\mathcal{RS} = \emptyset$;
  for $i = 1, \ldots, N_D$ do
    foreach $\tilde{x}_i \in$ random_sample(unfold($\mathcal{UC}_i$)) do
      $\mathcal{F} = \text{generate_meta_feature}(\tilde{x}_i)$;
      // use $h'$ to infer the most likely activity label $a$ and its probability $pr$;
      $(\tilde{x}_i, a, pr) = h'(\mathcal{F})$;
      $\mathcal{RS} = \mathcal{RS} \cup \{(\tilde{x}_i, a, pr)\}$
    end
  end
  // rank the result $\mathcal{RS}$ by the inferred probabilities in a descending order and select the top $k$ instances to update the corresponding base learners and ensemble learner
  $T_k = \text{select(rank(\mathcal{RS}), k)}$;
  // use Definition 5 to remap the predicted labels of instances in $T_k$ to the closest labels in each of their own datasets
  remap_labels($T_k$);
  for $i = 1, \ldots, N_D$ do
    // update the labelled and unlabelled clusters in each dataset
    add($\mathcal{LC}_i, T_k$);
    remove($\mathcal{UC}_i, T_k$);
    // update the base learner $h_i$ with the instances in the $T_k$ that belong to the $i$th dataset
    $h_i = \text{update}(h_i, T_k)$;
    update_model_accuracy($h_i$);
  end
  $h' = \text{update}(h', T_k)$
end

(1) how much training data are needed to achieve reasonably good accuracies? We start with the smallest training example 2, and gradually increase it up to 100 with a step size 2. We will monitor the increase of recognition accuracies over the increase of training data.

(2) Will it perform better than the state-of-the-art approaches? We compare the XLearn algorithm with baseline classification, cotraining and active learning algorithms.

(3) To what extent of heterogeneity can the XLearn approach perform robust cross learning? We will compare the performance of different combinations of datasets in XLearn and uncover how heterogeneity impacts the effectiveness of cross learning. This will serve as a guidance for selecting the most appropriate datasets to perform cross learning.

(4) Will XLearn enable more targeted annotations so as to reduce the annotation to an extreme while still achieving high accuracies? We will demonstrate a selection strategy to choose examples for annotation so that the performance of XLearn can be optimised. This will help to design a more practical way to using XLearn.

Datasets. We evaluate XLearn on four publicly available, real-world datasets that capture typical activities and that, more importantly represent common types of smart home datasets in terms of the number of sensors and residents, different spatial layouts, and the degree of inherent noise. We deliberately select these datasets that exhibit different level of heterogeneity, which can provide us a comprehensive view of the effectiveness of the proposed technique.
The first three datasets\(^2\) (denoted as House A, B, and C respectively in the following) are collected by the University of Amsterdam from three real-world, single-resident houses which were instrumented with wireless sensor networks \(^{[17]}\). These three datasets record 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 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. Even though the activity sets are the same from these datasets, they are recorded from three different real-world, residential settings that have different spatial layouts and host different subjects.

The fourth dataset\(^3\) (denoted as House W) is the interleaved activities of daily living from the CASAS smart home project \(^{[7]}\). This dataset was collected in a smart apartment testbed hosted at Washington State University during the 2009-2010 academic year. The apartment was instrumented with various types of sensors to detect user movements, interaction with selected items, the states of doors and lights, consumption of water and electrical energy, and temperature, resulting in 2.8M sensor events. The apartment housed two people, R1 and R2, who performed their normal daily activities during the collection period, including working, sleeping, or making meals. Our experiments consider the following 13 activities: R1_Sleep, R1_Work, R1_Wander in room, R2_Sleep, R2_Work, R2_Wander in room, Meal_Preparation, Watch_TV, Personal_Hygiene, Bathing, Leave/Enter_home, Housekeeping, and Eating. That is, we have merged some of the activity labels in the common areas together; for example, instances that are labelled R1_Meal_Preparation and R2_Meal_Preparation are changed to a common label Meal_Preparation. The reason is that XLearn itself is not able to distinguish the users when they perform the same type of activities in a common area. This can be done after plugging another machine learning technique \(^{[10]}\) after XLearn that is dedicated to learn fine difference between users. Figure 5 presents the similarity between activities in House W and the other three houses, which is calculated from Definition 4. As we can see, even with the simplification on House W, there exists relative large heterogeneity in these houses’ activity sets. There is no clear one-to-one mapping between the activities from these two datasets, and some activity like Sleep in House A can equally map to 6 activities in House W. All these issues make sharing and learning the labels on each dataset difficult.

\(^{2}\)https://sites.google.com/site/tim0306/datasets

\(^{3}\)http://casas.wsu.edu/datasets/
We have used Definition 5 to remap an inferred activity label to the most similar activity to the
expected activity label. When the sizes of the datasets are imbalanced – i.e., activities out of interest. After filtering, we segment sensor events into
a 60-second interval [16] and extract sensor features as defined in Definition 1. In the end, the total number of instances for House A, B, C, and W are 505, 497, 474, and 70053.

**Methodology.** The experiment setup is described as follows. The training data consists of \( n \times N_D \) labelled examples where \( n \) instances are are randomly selected from each of the \( N_D \) datasets and the test data will be all the remaining examples in each dataset. For example, if there are 3 datasets containing of 100, 200 and 300 examples respectively, given \( n = 4 \), then the training set will be composed of 12 labelled examples from these 3 datasets and the test set will include 588 examples altogether. In the prediction process, XLearn assigns the most confident label to each test example, and the label can come from a dataset different from the test example. We iterate \( n \) from 2 to 100 with a step size of 2. For each \( n \), we run 100 iterations to balance the differences in randomly generated annotated data. We choose the number over the percentage for training data because we would like to examine the extreme situations where the training examples are as small as possible.

When the sizes of the datasets are imbalanced – i.e., when one dataset contains millions of sensor events while another only contains thousands – it is hard to choose the smallest percentage.

**Metrics.** We examine the following accuracy metrics: (1) **Overall accuracy** \( O \), which measures the ratio of the learnt labels being the same as the true labels and is a commonly-used metric for activity recognition. However, the datasets of use have imbalanced activity distributions, so the overall accuracy can be misleading when the classifier that is optimised for this metric will always select the majority class while ignoring the minority class. (2) **Macro F1 score** \( F1 \), which is averaged F1 scores across all the activity classes and often used in the face of skewed class distribution. (3) **Similarity score** \( S \), which measures how close a learnt label is to a true label. In XLearn, the learnt labels can come from other datasets which might not share the same collection of activity labels. We have used Definition 5 to remap an inferred activity label to the most similar activity to the corresponding dataset, but the exact match can be difficult.

**Parameter and Algorithm Configuration.**

**Clustering algorithms.** We have employed the two state-of-the-art clustering algorithms: DBSCAN and KMEANS++. There is no real significance to these choices: a novel clustering algorithm or a sophisticated distance metric is not the main objective or contribution of this work. We assume that a more effective clustering algorithm could lead to a better performance.

To determine the optimal number of clusters, we use the F-test on the KMEANS++ algorithm; we set \( k \) from the number of activity labels that have occurred in the training data to \( \sqrt{n} \) (\( n \) being number of patterns in a cluster), and for each \( k \) we calculate the percentage of explained variance, which is the ratio of the between-cluster variance over the overall variance. An abrupt change in the percentage of explained variance suggests the corresponding \( k \) is an optimal solution.

We use the mutual information validation technique on the DBSCAN algorithm, based on information entropy and measures how likely it is that we reduce the uncertainty about the clustering \( C \) of a random element when knowing its cluster in another clustering \( C' \) of the same set of elements. We start with the mean distance between any two sensor features within the training set. We then increase it by a standard deviation of distance measures within each activity. For each setting, we will run the DBSCAN algorithm, and compute the mutual information score until we find the optimal result.

Classification. To choose a base learner for each dataset, we want a simple classifier with good prediction performance for a small number of training data. To find such a classifier we have explored a collection of the state-of-the-art classification algorithms: Logistic Regression, Naïve Bayes, Random Forest, J48 Decision Tree, Support Vector Machine, and Neural Networks. None of them have performed significantly better than the others. In the end, we use Naïve Bayes for the base learner algorithm for two reasons: the technique takes the least time to train, and generates quite small models. For example, on the House W dataset trained with 8 training examples, the Naïve Bayes classifier is about 70K, while the random forest is about 100MB. The small size of the classifier can make it easier to deploy on resource-constrained devices. Also the XLearn algorithm is mainly used at the early stage of the system to quickly acquire labels from other datasets. As the labelled examples accumulate, we can always replace the simple classifier with a more powerful one – although it is not clear this would result in better final classification.

Often an ensemble classifier is a logistic regressor, and we have compared results with different combinations of the above classifiers. Again there is no significant difference in performance in these combinations.

Ensemble labelling. In Algorithm 5, we randomly sample a subset of unlabelled clusters for updating. The sampling rate is set up as 50%. In XLearn we will need to choose top k examples for the next training iteration. We have experimented with different numbers from 5 up to 100 with a step size 5. The smaller k, the better the accuracies. However, a smaller k generates a lot of iterations and takes long time to converge. For the sake of performance, we choose k to be 20, which can achieve good accuracies with acceptable run times.

6 RESULTS AND DISCUSSION
In this section, we will present and discuss evaluation results.

6.1 Cluster labelling
Figure 6 presents the ratio and F1 scores of the cluster labelling process with the DBSCAN and KMEANS++ algorithms on all the datasets. We consider the label expansion process is effective in that it achieves high accuracies.

The theoretically best F1 we can achieve is $F_1^{best} = \min\left(\frac{N_c}{N_c}, \frac{N_{ct}}{N_c}\right)$, where $N_c$ is the total number of activity classes in a dataset, and $N_{ct}$ is the number of activity classes being labelled in the training data. It assumes that on each class $i$, $F_1^i$ achieves 100% of precision and recall. In the extreme situation, when there exist only 2 training instances from 2 different activity classes, then the $F_1^{best}$ will be 28% on House A, B, and C with $N_c = 7$, and 15% on House W with $N_c = 13$. As we can see in Figure 6, the F1 scores are above 20% on the House A, B, and C, and above 10% on the house W. Therefore, we conclude the cluster labelling has successfully recognised observed activities on each dataset. The results also show that there is no significant difference between DBSCAN and KMEANS++ in labelling accuracies. For the sake of performance, in the following experiments, we will use only with the KMEANS++ algorithm that runs much faster than DBSCAN.

6.1.1 Effectiveness of semantic cosine. To demonstrate the effectiveness of the semantic cosine metric, we compare the ratio and F1 scores of using the semantic cosine and cosine as the distance metric in the KMEANS++ algorithm. Results in Figure 7 show that semantics cosine outperforms cosine in the label expansion process. These distance metrics achieve similar labelling ratios, suggesting that both have a similar capacity of clustering close instances together. However, semantic cosine has achieved higher accuracies than cosine, indicating that the knowledge on sensor similarity does help to group sensor features that might belong to the same activities. With
We can see much better improvement for the datasets with a more diverse set of sensors, especially the wider coverage of sensor features, the more effective of finding the similar features. The increase in training data, the improvement in accuracies is becoming more and more significant. The wider coverage of sensor features, the more effective of finding the similar features.

The label expansion process can effectively label semantically similar examples. From the label expansion and are presented in Figure 8. It shows that with the expanded training set, the classifier can achieve better accuracies in all datasets. We can see much better improvement for the datasets with a more diverse set of sensors, especially the House W. The reason is that when the original training set is unable to capture all the possible combinations of sensor activations for a certain activity, semantic clustering helps to expand the training set with the instances that include semantically similar sensors.

### 6.1.2 Effectiveness of expanded training set
To evaluate the effectiveness of the expanded training set, we compare the recognition accuracies by applying Naïve Bayes on both the original and expanded training set. The accuracies are measured in O and F1 and are presented in Figure 8. It shows that with the expanded training set, the classifier can achieve better accuracies in all datasets. We can see much better improvement for the datasets with a more diverse set of sensors, especially the House W. The reason is that when the original training set is unable to capture all the possible combinations of sensor activations for a certain activity, semantic clustering helps to expand the training set with the instances that include semantically similar sensors.

### 6.2 Ensemble Labelling
Figure 9 presents the overall accuracies and F1 scores of XLearn, baseline learning, cotraining, and active learning classifiers on each dataset. Here baseline learning means that we train Naïve Bayes
with the original training set and test on the remaining data. With active learning we use Naïve Bayes with the information entropy uncertainty sampling strategy. We implement the cotraining algorithm from the original paper [3]; that is, we order the test data with their prediction confidence in a descending order, choose a top-$p$ percentage of the test data to update the classifier, and iterate the process until all the test data are labelled. We run $p$ from 10% to 50% with a step 10%. Because the results are similar, we only report the results on 10% and leave the rest accessible online.

The results in Figure 9 show that base classifiers and cotraining perform similarly and worst among all. The main reason is that both approaches suffer from a very small number of training data. Active learning produces higher overall accuracies and F1 scores when the number of training data is small, as it allows to acquire labels on uncertain examples from users. We report the query ratio – the percentage of test examples is queried to gain the labels. As we can see, when the number of training data is only 2, the query ratio can be between 10% and over 20%, which means 50 or 100 examples being queried, given that each of the House A, B, and C datasets contains about 500 examples in total. This makes the actual amount of training data is much higher than the training data that are used in the other two algorithms. This also explains both $O$ and $F1$ of active learning do not change much with the increase of training data.

The other reason that active learning performs better at the early stage is that it 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 XLearn 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

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4All the XLearn results can be accessed at https://drive.google.com/open?id=1tQkB0ERk74sTmFi5aNkDCrecPdngEC67P.
Fig. 9. XLearn outperforms baseline, cotraining and active learning algorithms because of the always availability assumption.

As presented in Figure 9, after a certain number of training data XLearn performs better than active learning, especially on the House B, C and W datasets. After a careful comparison of the
results, we identify two reasons. First of all, all these three datasets are more noisy than House A, which makes it difficult to estimate proper uncertainty threshold and thus lead to ineffective active learning. Secondly, XLearn has a more complex learning workflow – a stacked ensemble taking input from base classifiers and clusters, which leads to higher accuracy than the active learning approach that only uses the base classifier.

XLearn achieves better F1 scores than both baseline and active learning classifiers. This suggests that sharing across different sets helps increase the number of activity classes; for example, we can acquire unobserved activity labels from the other datasets. It is less likely that all the datasets are only annotated by the same set of activities.

6.3 Impact of Heterogeneity

As we claim XLearn works on heterogeneous datasets, we deliberately select these diverse datasets to demonstrate to what extent of heterogeneity datasets can be to achieve effective sharing. Here we study the impact of different combinations of datasets on XLearn’s performance, which will shed light on the sharing strategies – how to select most appropriate datasets to share, if there exists a large number of datasets. Figure 10 presents XLearn’s accuracies $O$, $F_1$, and $S$ on House A and W with four different combinations. On House A in Figure 10, when the amount of training data is small, the more data being available (e.g., ABC vs. AB) and the closer the datasets (e.g., AB vs. AW), the better accuracy is achieved. On House W, House B shares more similarity with House W given that the number of sensors in House B is larger, so we can observe higher accuracy on House W when House B is involved. When the training data accumulates, the difference between the combinations gets smaller, as each dataset will start using their own labels, which demonstrates XLearn’s ability to preserve activity models for individuals to prevent negative transfer. We have experimented all the other different combinations of datasets to further evaluate the impact of heterogeneity on the effectiveness of transfer learning, and the trend is similar.

The other observation is higher similarity scores of learnt activity labels with true labels, compared to overall and class accuracies. Looking at the inference results, we find that this mainly happens with the activities share a similar collection of objects and locations; e.g., ‘breakfast’, ‘dinner’, or ‘meal preparation’. As we are not using temporal features in the current experiments, these activities are difficult to select with the coarse-grained activity similarity in Definition 5. For example, if a learnt label on an instance on House A is ‘meal preparation’ from House W, then remapping this activity label to ‘breakfast’, ‘dinner’, and ‘drink’ can be less accurate. The reasons that we do not use temporal features are two folds. Firstly, in this work we want to focus on sensor signatures to understand basic sensor and activity mapping. Secondly, temporal features
can be specific to individuals or the way of annotating activities, which we consider can be an extra
application-level rule on top of XLearn.

Compared to our previous work SLearn [37], the sharing data algorithm with Naive Bayes on
SLearn outperforms XLearn when the training samples are really small; i.e., when 2 or 4 labelled
examples are selected from House A, B and C, SLearn with NB on House A can achieve the overall
accuracy of 68% and the macro F1-score of 50%. The reason is that gathering all the training
data together will improve the effectiveness of learning, while building a non-trivial stacked
ensemble will consume more training data than a simple classifier like Naive Bayes. However,
XLearn achieves much better accuracies when the training data contains slightly more examples;
i.e., 8. XLearn converges much faster than the sharing classifier algorithm on SLearn, and can enable
customised learning on each dataset compared to the sharing data algorithm on SLearn.

### 6.4 Strategic selection of examples to label

All the above examples are based on randomly sampled examples from the datasets for annotation.
What if we can select examples to label? Would such selection improve the recognition accuracies?
Driven by these questions, we run another set of experiments. For a given number of training data
$n$, we rank the clusters based on their size in a descending order and select the centroid examples
of the top $n$ clusters to label. The intuition is to select most representative examples to label from
each dataset, which is one common strategy in active learning to select informative instances
to annotate. As we can see in Figure 11, we can achieve much improved accuracies across all
the datasets. The overall and class accuracies can be as high as nearly 70% and 40% when only 2
examples from each dataset are selected for training. This is largely because selecting the most
frequent and representative examples from each dataset collectively increase the performance.

![Select examples to label](image)

**Fig. 11.** Overall accuracies and F1 scores with selected examples to label. With XLearn, we can effectively
target examples to annotate, leading to much higher recognition accuracies than random annotations. This
will help to further reduce the amount of training data.

### 6.5 Limitations and future work

The current work opens a new research direction towards a sustainable and scalable activity
recognition; that is, how to address the annotation challenge by taking advantage of the scalability
– leveraging data from a large number of users. The current solution is only a starting point and it
can be improved in the following aspects.

#### 6.5.1 Selection of datasets to share

If there exist a number of datasets, we can select the datasets
which will not only lead to higher recognition accuracies but also reduce the computation complexity.
Those we select can be based on the similarity between their sensors and the similarity between
their activities. Another direction that can usefully be pursued is to select users who have similar
activity routine as the target users. These routine can be either learnt through their reported activities, or through the other information like age, occupation, and family structure.

6.5.2 Strategic selection of examples. We have demonstrated the effectiveness of selecting examples to label on XLearn’s performance. We only experiment with the most intuitive solution – selecting the most representative activities. However, the most representative activities selected from each dataset can be similar; i.e, the majority classes. In the future, we can take into account of diversity: that is, how to cover a set of activity classes as large as possible. For example, we can select future examples that are different from the already-labelled examples in terms of their semantic difference and temporal features. Promoting diversity in selection of examples can further reduce the number of training data acquired from human operators.

6.5.3 Knowledge engineering. The generality and feasibility of the smart home ontologies have been demonstrated across different smart home environments without any modifications [39]. To perform feature space remapping, we need to take the sensor deployment file to map sensors to their corresponding location and object concepts. This limits the application of this knowledge-driven approach to a certain extent in that our approach works better in a setting where sensors are more or less fixed deployed and the semantic mapping between a source and target environment is achievable, rather than an open environment where each sensor is mobile and can join and leave the environment at any time. For example, when sensors are removed or moved, or a new sensor is introduced, we will need to remap sensors and this effort is unavoidable in our current design. Also to enable activity space remapping, we need to take a collection of pre-defined activities from each dataset to compute the activity similarity matrix. With a general methodology of human activity recognition [40], the system designer pre-defines a closed set of activities of interest, which are often the requirements from applications such as personal healthcare. Then driven by the activities, the designer will select a range of ambient and/or wearable sensors that can potentially detect these activities. The designers will then start collecting sensor data for a short period of time and annotate them with activity labels. With the annotated data, the developers will train a computational model for activity recognition.

In XLearn, we take the pre-defined activity description from each dataset to prepare for activity label remapping. What XLearn aims to achieve is to reduce the time and effort on annotating sensor data with activity labels. The pre-defined activity collection can evolve over time; that is, the system designers might add new activities to monitor. Then XLearn will take the new description to update the models and share the activity across the other datasets. The activity label remapping can incur extra knowledge engineering effort from the designers of smart home systems. However, when different environments use different activity labels, there exists few techniques to easily build mapping between them.

7 CONCLUSION

This paper proposes the XLearn algorithm to significantly reduce annotation burdens on individual users by distributing that burden across multiple users. We have shown that such distribution can result in more robust activity recognition by covering a diverse set of activities and activity patterns. It can shorten the initialisation phase of an activity recognition system by working on the labelled examples collected simultaneously across multiple users. In addition, the lighter burden of annotation potentially supports long-term sustainable activity recognition that only requires users to sporadically report their new activities over the deployment of an activity recognition system. More specifically, XLearn:
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- supports robust activity recognition that only requires limited number of training data (for example with about 6 annotated examples from each dataset, we can achieve nearly 85% overall accuracies and over 75% F1 scores);
- supports sustainable activity recognition in the sense that as long as each user can contribute a few annotated examples over time, we can share and transfer learning across different users; and
- is scalable in that it takes advantage of the large number of users: the more users contribute their annotations, the better we can learn and the less annotations we expect from individuals.

With these very promising results, we can envision another way of performing activity recognition: we can start recognition without the need for an explicit training period. When the sensors are deployed, we can start collecting sensor data and users’ annotations over time at their own pace. If one user has annotated a few examples of ‘preparing breakfast’ and the user in another environment has annotated a few examples of ‘taking bath’, then XLearn can start recognising both activities for both users. This will support long-term incremental activity recognition in that it can accommodate new activities annotated by different users over time.

The current design of XLearn relies on the ontologies to map feature spaces from different sensorised environments, which facilitates bridging their binary sensor feature space. To assess the generality of the approach, we will adapt XLearn to the other types of sensor data. To do so, we might need to replace the ontologies with other feature space remapping techniques.

Also XLearn mainly targets at meaningful activities; that is, the activities can be semantically described. This leads to another limitation of XLearn in detecting miscellaneous or ‘other’ activities.

In the future, we will look into how to combine XLearn with other techniques, such as uncertainty estimation strategies in active learning or new activity discovery [11], to recognise ‘other’ activities, as the ones that cannot be confidently classified as any meaningful activity.

REFERENCES


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