FlashProfile: Interactive Synthesis of Syntactic Profiles

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ABSTRACT

We address the problem of learning comprehensive syntactic profiles for a set of strings. Real-world datasets, typically curated from multiple sources, often contain data in various formats. Thus any data processing task is preceded by the critical step of data format identification. However, manual inspection of data to identify various formats is infeasible in standard big-data scenarios.

We present a technique for generating comprehensive syntactic profiles in terms of user-defined patterns that also allows for interactive refinement. We define a syntactic profile as a set of succinct patterns that describe the entire dataset. Our approach efficiently learns such profiles, and allows refinement by exposing a desired number of patterns.

Our implementation, FlashProfile, shows a median profiling time of 0.7 s over 142 tasks on 74 real datasets. We also show that access to the generated data profiles allow for more accurate synthesis of programs, using fewer examples in programming-by-example workflows.

1. INTRODUCTION

In modern data science, most real-life datasets lack high-quality metadata — they are often incomplete, erroneous, and unstructured [15]. This severely impedes data analysis, even for domain experts. For instance, a merely preliminary task of data wrangling (importing, cleaning, and reshaping data) consumes 50–80% of the total analysis time [33]. Prior studies show that high-quality metadata not only help users clean, understand, transform, and reason over data, but also enable advanced applications, such as query optimization, schema matching, and reverse engineering [47, 38]. Traditionally, data scientists manually inspect samples of data, or create aggregation queries. Naturally, this approach does not scale to modern large-scale datasets [38].

Data profiling is the process of automatically discovering useful metadata (typically as a succinct summary) for the data [6]. In this work, we focus on syntactic profiling, i.e.

\[ \text{Learn meaningful succinct descriptions from data.} \]

1 Work done during an internship at Microsoft.

| Birth Year | Percentage Data Coverage | Desired |
|------------|--------------------------|---------|
| 1900       | 100                      |         |
| 1877       | 97                       | 5       |
| 1866       | 89                       | 10      |
| 1893       | 82                       | 5       |
| 1888       | 18                       | 2       |
| 1872       | 30                       | 10      |

(a) Dataset

(b) Suggested and refined profiles

Figure 1: Profiles generated by FlashProfile for a real dataset. Atoms are concatenated with \( \cdot \), and superscripts indicate their repetitions. The number of matches for each pattern is shown on the right.

learning a succinct structural description of the data. We present FlashProfile, a novel technique for learning syntactic profiles that satisfy the following three desirable properties:

Comprehensive: We expose the syntactic profile as a set of patterns, which cover 100% of the data.

Refinable: Users can interactively refine the granularity of profiles by requesting the desired number of patterns.

Extensible: Each pattern is a sequence of atomic patterns, or atoms. Our pattern learner, \( Z_L \), includes a default set of atoms (e.g., digits and identifiers), and users can extend it with appropriate domain-specific atoms for their datasets.

Example: Figure 1 shows the syntactic profiles generated by FlashProfile for an incomplete and inconsistent dataset containing birth years. The profiles expose rare patterns in the data that are otherwise hard to notice. For example, the automatically suggested profile in Figure 1(b) reveals dirty years of the form \( 18\times \) Digit \( \times \) Digital
Similarly, *Potter’s Wheel* only detects the most frequent pattern in the dataset. Furthermore, for our example from Figure 1(a), it detects the most frequent pattern as `int` but not its fixed length 4. We give a detailed comparison in §8.

**Our Technique:** We profile a given dataset by first partitioning it into syntactically similar clusters of strings, and subsequently learning a succinct pattern describing each cluster. To facilitate user-driven refinement of the results into more clusters, we construct a hierarchical clustering over the dataset. This enables efficient extraction of clusters with desired granularity by splitting the hierarchy at an appropriate height. Two major challenges to constructing the hierarchy are — (1) defining an appropriate dissimilarity measure that that allows domain-specific profiling, and (2) computing all pairwise dissimilarities, which is typical for hierarchical clustering, is expensive for large datasets.

Our key insight towards addressing the first challenge is that, the desired measure of dissimilarity is not a property of the strings per se, but of the patterns describing them over the user-defined language. We define syntactic dissimilarity based on costs of patterns — a low cost pattern describing two strings indicates a high degree of syntactic similarity. To address the second challenge, we show sampling and approximation techniques which reuse previously learned patterns to approximate unknown pairwise dissimilarities. This enables hierarchical clustering using very few pairwise dissimilarity computations. In essence, we present a general framework for profiling, based on an efficient hierarchical clustering technique which is parameterized by a pattern learner $L$, and a cost function $C$ over patterns.

**Implementation and Evaluation:** Our implementation, FlashProfile, uses a pattern learner based on *inductive program synthesis* [32] — an approach for learning programs over an underlying domain-specific language from an incomplete specification (such as input/output examples). We formally define the synthesis problem for our pattern language $L_P$, and present (1) a *sound and complete* pattern learner $L_P$ over a user-specified set of atoms, and (2) a cost function $C_P$ over $L_P$ patterns. We have implemented $L_P$ using PROSE [4] (also called FlashMeta [40]), a state-of-the-art inductive synthesis framework.

We evaluate our technique on 74 publicly-available real datasets\(^1\) collected from online sources. Over 142 tasks, FlashProfile achieves a median profiling time of 0.7s, 77% of which complete in under 2s. Apart from being refutable interactively, we show that profiles generated by FlashProfile are more expressive compared to three state-of-the-art existing tools, owing to its extensible language.

**Applications:** The benefits of comprehensive profiles extend beyond data understanding. An emerging technology, programming by examples [32, 19, 21] (PBE), provides end users with powerful semi-automated alternatives to manual data wrangling. A key challenge to its success is finding a representative set of examples which best discriminate the desired program from a large space of possible programs [37]. We show that FlashProfile helps existing PBE systems by identifying syntactically diverse inputs.

We have investigated 163 scenarios where Flash Fill [18], a popular PBE system for string transformations, requires > 1 example to learn the desired transformation. In 84% of them, the representative examples belong to different syntactic clusters identified by FlashProfile. Moreover, for 86% of them, an interaction guided by the profile completes the task in a *minimum possible* number of examples.

In summary, we present the following major contributions:

- (§ 3) We define interactive profiling as a problem of hierarchical clustering based on syntactical similarity, followed by qualifying each cluster with a pattern.
- (§ 4) We propose a novel dissimilarity measure which is superior to traditional string-similarity measures for estimating syntactic similarity. We also present sampling and approximation strategies for efficiently constructing hierarchies using the proposed measure.
- (§ 5) We instantiate our technique as FlashProfile, using program synthesis for learning patterns over the language $L_P$. Our learner $L_P$ also supports user-defined patterns.
- (§ 6) We evaluate FlashProfile’s performance and accuracy across 142 tasks on real-life datasets, and compare profiles generated by FlashProfile to state-of-the-art tools.
- (§ 7) We show how FlashProfile helps PBE systems by identifying a representative set of examples for the data.

2. **MOTIVATING SCENARIO**

In this section, we discuss a practical data analysis task and show the benefit of comprehensive data profiles. Consider the task of gathering descriptive statistics (e.g. range and mode) of the data in Figure 1(a). The following Python script is a reasonable first attempt after a quick glance on the data:

```python
# collect clean entries
clean = [int(e) for e in years if e != '?']
min_year = min(clean)
max_year = max(clean)
mode_year, mode_count = Counter(clean).most_common(1)[0]
```

But it fails when `int(e)` raises an exception on e == `'`. Thus, the user updates the script to clean the data further:

```python
# try again
clean = [int(e) for e in years if len(e) == 4]
```

Having encountered only 4-digit years and ? in the first sample, she simplifies the script to consider only clean 4-digit years. Now the script runs to its completion and returns:

```python
min_year = 1813
max_year = 1875
mode_year = 1903
mode_count = 118
```

However, the `mode_count` value is incorrect because the analysis script ignores any approximate entries with a trailing ? (e.g., 1875?). Since the user is unaware of such entries after a quick glance on the data, she does not realize that the computed value is incorrect.

A more cautious analyst might have discovered entries like 1875? after adding a second check to filter out the missing entries. However, such trial-and-error pattern discovery requires several time-consuming iterations to cover the entire dataset. The analyst also needs to manually inspect the failing cases and find a general pattern to handle them properly. Moreover, this approach works only if she knows how to validate the data type (note the `int(e)` in `clean`), which might be difficult for non-standard types.

**Wrangling by Examples:** Instead of manually analyzing the data, the user may choose a semi-automated technique like Flash Fill. She loads her data in Excel and provides a few examples, expecting Flash Fill to learn the transformation for cleaning any dirty years by extracting their integer parts. Typically users provide examples for the first few rows. Flash Fill then synthesizes the *simplest generalization* [18] over the provided examples.

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\(^1\) Available at: https://github.com/SawatPahdi/ProfilingExperiments
In this case, Flash Fill produces identity transform since the first several inputs are all clean 4-digit years. Fixing this mistake requires manual inspection, which reveals a discrepancy only at the 84th row. Furthermore, if the user invokes the MIN, MAX, or MODE functions on the output column, Excel silently ignores all non-numeric entries, thus producing incorrect results without any warnings.

As we show in §7, a profile of the input column allows Flash Fill to learn the desired transformation with fewer examples. Instead of simply taking examples in the order they appear in the dataset, Flash Fill proactively requests it on the most discrepant row w.r.t. the previous examples.

3. OVERVIEW

In this section, we informally explain our technique, and its instantiation as FlashProfile. The goal of syntactic profiling is to learn patterns summarizing the given data, which are neither too specific nor too general (to be practically useful). Here, the dataset itself is a trivial overly specific profile, whereas the regex `.*` is an overly general one.

**Definition 3.1, Syntactic Profile:** Given a set of strings \( S \), syntactic profiling involves learning (1) a partitioning \( \{S_1, \ldots, S_k\} \) of the set \( S \) and (2) a set of syntactic patterns \( P = \{P_1, \ldots, P_k\} \), such that each \( P_i \) succinctly describes the partition \( S_i \). The set \( P \) is called a syntactic profile of \( S \).

At a high level, our approach uses a clustering algorithm, which invokes pattern learning on demand. In Figure 2, we outline our Profile algorithm, which is parameterized by:

- a pattern learner \( \mathcal{L} \) that accepts a set of strings \( S \), and returns a set of patterns consistent with them.
- a cost function that returns a numeric cost for a given pattern with respect to a given dataset.

Profile accepts a dataset \( S \), with bounds \([m, M]\) for the number of patterns desired, a real number \( \theta \) (a sampling factor), and returns a profile \( \hat{P} = \{\langle S_1, P_1 \rangle, \ldots, \langle S_k, P_k \rangle\} \) with \( m \leq k \leq M \). First, it invokes \textsc{BuildHierarchy} to construct a hierarchy \( H \) (line 1), which is subsequently partitioned by \textsc{Split} (line 3). Finally, for each partition \( X \), \textsc{LearnBestPattern} is invoked (line 4) to learn a pattern \( P \), which is added to \( \hat{P} \) paired with its partition \( X \) (line 5).

3.1 Hierarchical Clustering with Sampling

\textsc{BuildHierarchy} yields a hierarchy (or dendrogram) over a given dataset \( S \), representing a nested grouping of strings based on their similarity [25]. For instance,
large number of consistent patterns. Using the learner $L$ and cost function $C$, we define the dissimilarity measure $\eta$ as the minimum cost incurred by any pattern for describing a given pair of strings. We evaluate $\eta$ in §6.1, and show its superiority over classic character-based similarity measures, and machine-learned regression models. Figure 3(b) shows dissimilarities incurred by FlashProfile for a few pairs of strings from our running example. We also show the least-cost patterns describing them using the default atoms listed in Table 1. The void (⊥) indicates a pattern-learning failure that arises because no pattern in FlashProfile matches both empty and non-empty strings. We associate equal measures of dissimilarity with ⊥ and Any*.

Adaptive Sampling and Approximation: While $\eta$ captures a high-quality syntactic similarity, it is expensive to compute. With it, each pairwise dissimilarity computation during AHC would require learning and scoring of patterns, making our technique too expensive for large real datasets.

We address this challenge by using a two-stage sampling technique. (1) At the top-level, FlashProfile employs a Sample→Profile→Filter cycle: we sample a small subset of the data, profile it, and filter out data that is explained by the profile learned so far. In §6.2, we show that a small sample of strings is often sufficient to learn a general pattern describing them. (2) While profiling each sample, our BuildHierarchy algorithm adaptively samples a few pairwise dissimilarities, and approximates the rest. The key domain-specific observation that enables this is, computing the dissimilarity for a pair of strings gives us more than just a measure – we also learn a pattern. We test this pattern on other pairs, to learn their approximate dissimilarity. This is typically much faster than learning new patterns.

Example 2. The pattern “1” $\circ$ Digit$^*$ learned for the strings {1817, 1901}, also describes {1872, 1901}, and may be used to compute their dissimilarity, without pattern learning. Figure 3(b) shows other such recurring patterns.

Although the pattern “1” Any$^+$ learned for {1901, 1872, 1875} also describes {1872, 1875}, there exists another pattern “187” Any$^+$ which indicates a lower syntactic dissimilarity (due to a larger overlap) between them. Therefore, the dissimilarities (and equivalently, patterns) to be sampled, need to be chosen carefully for accurate approximations. We detail our adaptive sampling, and approximation algorithms in §4.2 and §4.3 respectively.

3.2 Pattern Learning via Program Synthesis

An important aspect of our clustering based approach to profiling, described in §3.1, is its generality. It may leverage any learning technique which provides:

- a pattern learner $L$ over strings, for:
  - computing dissimilarity, and approximations
  - exposing the profile to end users

Our language $L$ describes a string $s$, i.e. $P[s]$ = True iff $s$ satisfies every constraint imposed by $P$. Patterns are defined as arbitrary sequences of atomic patterns (atoms). We assume a default set of atoms (listed in Table 1) which may be extended with arbitrary pattern-matching logic. A pattern describes a string, if each successive atomic match succeeds, and they match the string in its entirety.

Example 3. The pattern “18” $\circ$ Digit$^* 2$ $\circ$ “?”, matches 1875, but not 1872 since the atom “?” fails to find a match. Unlike regexes, patterns in $L_P$ must match entire strings. For instance, “18” $\circ$ Digit$^* 2$ matches 1872, but not 1875 since the suffix ? is left unmatched.

Pattern Synthesis: The inductive synthesis problem for pattern learning is: given a set of strings $\mathcal{S}$, learn a pattern $P \in L_P$ that describes every string in $\mathcal{S}$, i.e. $\forall s \in \mathcal{S}: P[s] = True$. Since $P$ is a sequence of atoms, we can decompose the synthesis problem for matching $P$ into smaller problems for matching individual atoms. However, a naïve approach of tokenizing each string to (exponentially many) sequences of atoms and computing their intersection is quite expensive. Instead, our learner $L_P: 2^\mathcal{S} \rightarrow 2^{L_P}$, finds the intersection in an incremental fashion, at the boundaries of atomic matches, by computing a set of compatible atoms. Furthermore, $L_P$ is sound and complete over $L_P$ patterns, restricted to the user-specified set of atoms.

$L_P$ is built using PROSE [4] (also called FlashMeta [40]), a state-of-the-art inductive synthesis framework that is being deployed industrially [41]. It performs a top-down walk over the specified DSL grammar, at each step reducing a given synthesis problem on the desired program to smaller synthesis problems on subexpressions of the program, based on the reduction logic specified by the DSL designer. We explain the details of $L_P$ in §5.2.
Cost of Patterns: Once a set of consistent patterns is identified, a variety of strategies may be employed to identify the most desirable pattern w.r.t. the dataset. Our cost function, \( C_P : L_P \times 2^S \rightarrow \mathbb{R} \), quantifies the suitability of each pattern with respect to the given dataset. Intuitively, it decides a trade-off between two opposing factors:

- **Specificity**: select a pattern that does not over-generalize
- **Simplicity**: select a compact (interpretable) pattern

**Example 4.** The pair \{Male, Female\} are matched by the patterns \( \text{Upper} \circ \text{Lower}^\dagger \) and \( \text{Upper} \circ \text{HexString} \circ \text{Lower}^+ \). Although the latter is a more specific pattern, it is overly complex. On the other hand, the pattern \( \text{Alpha}^+ \) is simpler and easier to interpret by end users, but is overly general.

To this end, each atom in \( L_P \) has an empirically fixed **static cost**, and a dataset-driven **dynamic weight**. The final cost of a pattern is the weighted sum of the cost of atoms that appear in it. We describe our cost function \( C_P \) in §5.3.

4. **Hierarchical Clustering**

In this section, we first explain our algorithm for building a hierarchy and learning a profile by sampling dissimilarities for only a few pairs of strings. In §4.4, we then discuss how to combine profiles generated over several samples of a dataset to generate an overall profile. Henceforth, we use the term **pair** to denote a pair of strings.

The procedure \textsc{BuildHierarchy}, listed in Figure 5, constructs a hierarchy \( H \) over a given dataset \( S \), assuming at most \( M \) clusters to be extracted, using the sampling factor \( \theta \). First, \( \textsc{SampleDissimilarities} \) computes a hashtable \( D \) (line 1), which maps only \( O(\theta M|S|) \) pairs sampled from \( S \), to the dissimilarities and best patterns computed for them. We formally define this dissimilarity measure in §4.1, and detail the \textsc{SampleDissimilarities} algorithm in §4.2. \( D \) is then used by \textsc{ApproxDMatrix} to create matrix \( A \) containing all pairwise dissimilarities over \( S \), approximating wherever necessary (line 2). Finally, \( A \) is used by \textsc{AHC} to construct the hierarchy \( H \), using the complete-linkage criterion \( \eta \) (line 3). We explain \textsc{AHC} based on approximations, in §4.3.

4.1 **Syntactic Dissimilarity**

We first formally define a syntactic dissimilarity measure over strings based on the patterns learned by \( \mathcal{L} \) (specifically, the one with least cost), which describe them together.

**Definition 4.1.** Syntactic Dissimilarity: The syntactic dissimilarity of identical strings is 0. For strings \( s_1 \neq s_2 \), we define the syntactic dissimilarity as the minimum cost incurred by any pattern to describe both:

\[
\eta(s_1, s_2) = \begin{cases} 
C_{\text{max}} & \text{if } V = 0 \\
\min_{P \in V} C(P, \{s_1, s_2\}) & \text{otherwise} 
\end{cases}
\]

where \( V = \mathcal{L}([s_1, s_2]) \) denotes the patterns learned for \( \{s_1, s_2\} \), and \( C_{\text{max}} \) denotes a high cost for a learning failure.

4.2 **Adaptive Measuring of Patterns**

Although \( \eta \) accurately measures the syntactic dissimilarity of strings over an arbitrary language of patterns, it does so at the cost of performance. Using \textsc{LearnBestPattern}, every pairwise dissimilarity computation requires pattern learning and scoring, which may be computationally expensive depending on the number of consistent patterns generated. Moreover, typical clustering algorithms [25] require all pairwise dissimilarities. To put this into perspective, even with a fast pattern learner requiring \( \sim 10 \) ms per dissimilarity computation\(^7\), one would spend over 3s on computing all 300 pairwise dissimilarities for profiling only 25 strings.

As shown in example 2, previously learned patterns may be used to estimate dissimilarities for other pairs. However, the patterns to be sampled must be chosen carefully. For instance, a pattern learned for a pair with very high dissimilarity may be too general, and may match many other pairs. But, approximations based on these matches would be inaccurate since the cost of this pattern may differ vastly from the least-cost patterns for other pairs.

Our adaptive sampling algorithm, shown in Figure 6, is inspired by the seeding technique of \textsc{k-means++} [8]. \textsc{SampleDissimilarities} accepts a dataset \( S \), an integer \( \tilde{M} \) that is provided by \textsc{BuildHierarchy} as \( \lceil \theta M \rceil \) (line 1, Figure 5), and samples \( O(\theta M|S|) \) pairwise similarities.

\(^7\) \textsc{FlashProfile}'s pattern learner \( \mathcal{L}_P \) has a median learning time of 7 ms per pairwise dissimilarity, over our benchmarks. For comparison, most recent synthesizers based on PROSE have a learning time of \( \sim 500 \) ms [31].
4.3 Approximately Correct AHC

Our approximation technique is grounded on the observation that testing whether a pattern matches a string, is typically much faster than learning new patterns. If the dissimilarity of a pair is unknown, we test the sampled set of patterns, and approximate the dissimilarity as the minimum cost incurred by any known pattern that is consistent with the pair.

Using the hashtable $D$ from SAMPLEDISSIMILARITIES, the method APPROXDMATRIX, listed in Figure 7, approximates a matrix $A$ of all pairwise dissimilarities over $S$. For identical strings, the dissimilarity is set to 0 (line 4). If the dissimilarity of a pair has already been sampled in $D$, we simply copy it to $A$ (line 5). Lines 7, 8 show the key approximation step that are executed if the dissimilarity of a pair $(s, s')$ has not been sampled. In line 7, we invoke TESTPATTERNS to select a set $X$ containing only those patterns $P$ from $D$, which describe both $s$ and $s'$, i.e. $P[s] = P[s'] = \text{True}$. If $X$ is non-empty, we simply select the least-cost pattern from $X$ (line 8). If $X$ turns out to be empty, i.e. no sampled pattern describes both $s$ and $s'$, then we invoke LEARNBESTPATTERN to compute $\tilde{\eta}(s, s')$ (line 10). We also add the newly learned pattern to $D$, in line 11, to use it for subsequent approximations.

Once we have all pairwise dissimilarities, we use a standard AHC algorithm [25], shown in Figure 8. Initially in line 1, each string starts in its own cluster, which form the leaf nodes in the hierarchy. In each iteration, we select the least dissimilar pair of clusters (line 3) and join them to a new internal node (line 4), till we are left with a single node (line 2), which becomes the root of the hierarchy. To select the least dissimilar pair of clusters, AHC algorithms require a linkage criterion (line 3). We use a classic complete-linkage criterion [48] over our pairwise dissimilarities:

$$\tilde{\eta}(X, Y | A) \triangleq \max_{s_1 \in X, s_2 \in Y} A[s_1, s_2]$$

where $A[s_1, s_2]$ indicates the dissimilarity of $s_1$ and $s_2$.

Compared to other linkage criteria, complete linkage has been shown to be more resistant to outliers and yield useful hierarchies in many practical applications [25].

After a hierarchy $H$ has been constructed, our PROFILE algorithm (listed in Figure 2) invokes the SPLIT method to extract $m \leq k \leq M$ clusters. If $m = M$, it simply splits the first $m$ levels (the top-most $m$ internal nodes) of the hierarchy. Otherwise, it uses a heuristic based on the elbow (also called knee) method. Between the $m$th and the $M$th level, it locates a level $k$ in the hierarchy after which the dissimilarities of subsequent levels do not seem to vary much.

4.4 Profiling Large Datasets

To scale our technique to large datasets, we implement a second round of sampling around our core PROFILE method. Recall that, the SAMPLEDISSIMILARITIES samples $O(\theta M | S)$ pairwise dissimilarities by selecting a set $\rho$ of most dissimilar strings, and computing their dissimilarity with all strings in dataset $S$. But, although $|\rho| = \Theta(M)$ is very small, $|S|$ is still very large (several thousands) for real-life datasets.

Our second sampling technique runs the PROFILE algorithm from Figure 2 on small chunks of the original dataset and combines the generated profiles. We observe that even a small randomly selected subset exhibits relatively frequent patterns in the dataset, and our learner $\mathcal{L}_p$ does not require more than a couple of strings to learn them in most cases.

We implement a simple Sample–PROFILE–Filter loop:

1. sample $|\mu M|$ strings from $S$,
2. add their profile to the current set of known patterns,
3. remove strings described by the known patterns from $S$.
In this section we describe the most similar partitions in the profile so far. We then merge the data in $X.Y$ to a single partition $Z$ (line 3), learn a pattern describing $Z$ (line 4), and update the profile $\tilde{P}$ by replacing $X.Y$ with $Z$ and its pattern (line 5). COMPRESSPROFILE repeats this entire process till the total number of patterns falls within the upper-bound $M$.

5. PATTERN SYNTHESIS

In this section we describe FlashProfile, which instantiates the proposed profiling technique based on clustering. We begin with a brief description of the language in §5.1. In §5.2, we present our pattern learner $\mathcal{L}_P$, which produces all patterns consistent with a given dataset using a user-specified set of atoms over $\mathcal{L}_P$. Finally, in §5.3, we provide a description of our cost function $C_P$ for $\mathcal{L}_P$ patterns.

5.1 The Pattern Language $\mathcal{L}_P$

Figure 10(a) shows formal syntax for our pattern language $\mathcal{L}_P$. Each pattern $P \in \mathcal{L}_P$ is a function $P : \text{String} \rightarrow \text{Bool}$ which embodies a set of constraints over strings. A pattern $P$ describes a given string $s$ i.e. $P[s] = \text{True}$, iff $s$ satisfies all constraints imposed by $P$. Patterns in $\mathcal{L}_P$ are defined in terms of atomic patterns:

Definition 5.1. Atomic Patterns (Atoms): An atom, $\alpha : \text{String} \rightarrow \text{Int}$ is an operator which given a string $s$, returns the length of the longest prefix of $s$ matched by it. Atoms only match non-empty strings, i.e. $\alpha(s) = 0$ indicates match failure for $\alpha$ on the string $s$.

$\mathcal{L}_P$ allows the following four kinds of atoms:

- Constant Strings: $\text{Const}$, only matches the string $s$. We use a string literal such as “data” to denote $\text{Const}$.data”.
- Regular Expressions: $\text{RegEx}$, matches the longest prefix matched by the regular expression $r$.
- t e c sets of characters
- $f$ functions String → Int
- $r$ regular expressions
- $s$ strings $\Sigma$
- $e$ non-negative integers

FlashProfile provides a default set of atoms listed in Table 1. However, the set of atoms may be extended by end users to generate rich profiles for their datasets.

Example 7. Atom Digit is $\text{Class}_2^D$ with $D = \{0, \ldots, 9\}$. Digit+ is $\text{Class}_2^D$, and Digit×2 is $\text{Class}_2^2$. The atom Digit×2 matches the string 04/23/2017 but not 2017/04/23. For the latter case, the longest prefix matched by Digit+ is 2017, of length 4 ≠ 2. However, Digit+ matches both strings, and returns 2 and 4 respectively.

Definition 5.2. Patterns: A pattern is defined by a sequence of zero or more atoms. The $\mathcal{L}_P$ expression $\text{Empty}$ denotes an empty sequence of atoms, which only matches the empty string $\varepsilon$. The pattern $\alpha_1 \circ \alpha_2 \circ \ldots \circ \alpha_n$ is denoted by $\text{Empty}(\text{SuffixAfter} (\ldots \text{SuffixAfter}(s_{\alpha_1}) : \ldots, s_{\alpha_k}))$ in $\mathcal{L}_P$, which matches a string $s_0$, iff $s_k = \varepsilon$ and $s_0 \in \mathbb{N}, \alpha_0(s_{i+1}) > 0$ where $s_{i+1} = s_i[\alpha_i(s_i): ]$ i.e. the remaining unmatched suffix of string $s_i$ after matching atom $\alpha_{i+1}$.

Formal semantics for atoms and patterns are shown in Figure 10(b). $s[t]$ denotes the $t^{th}$ character of $s$, and $s[t:j]$ denotes the substring of $s$ from the $t^{th}$ character, till before the $j^{th}$ character. $\alpha$ may be omitted to indicate a substring extending till the end of $s$. In $\mathcal{L}_P$, the $\text{SuffixAfter}(s, \alpha)$ operator computes $s[\alpha(s): ]$, or raises an error if $\alpha(s) = 0$.

Example 8. Consider the following URLs collected from a dataset containing flight data for various destinations:

- http://www.jetradar.com/flights/EsaAla-ESA/
- http://www.jetradar.com/flights/Mumbai-BOM/
- http://www.jetradar.com/flights/Bangalore-BLR/
- http://www.jetradar.com/flights/LaForges-YLF/

The following pattern describes these URLs:

“http://www.jetradar.com/flights/”

Upper α° “+” = Upper° “+”

\[ \text{Pattern } P[s] := \text{Empty}(s) \quad | \quad P[\text{SuffixAfter}(s, \alpha)] \]

Atom $\alpha := \text{Class}_2^D|\text{RegEx}, \quad | \quad \text{Const}$ (a) Syntax of an $\mathcal{L}_P$ pattern $P$.

Empty $(s) \downarrow \text{true}$

$s = s_0 \circ s_1$ $\quad (s_a) = |s_0| > 0$

$\text{SuffixAfter}(s, \alpha) \downarrow s_1$

$\text{Funct}(s) \downarrow f(s)$

$s'' = s \circ s''$

$\text{Const}(s) \downarrow |s|$

\[ \begin{align*}
\text{Empty}(s) & \downarrow \text{true} \\
\text{Funct}(s) & \downarrow f(s) \\
\text{Const}(s) & \downarrow |s|
\end{align*} \]

(b) Semantics of $\mathcal{L}_P$ patterns. The judgement $E \downarrow v$ denotes that the expression $E$ evaluates to a value $v$.

\[ \begin{align*}
\text{Empty}(s) & \downarrow \text{true} \\
\text{Funct}(s) & \downarrow f(s) \\
\text{Const}(s) & \downarrow |s|
\end{align*} \]
5.2 Synthesis of \( \mathcal{L}_P \) Patterns

FlashPro's pattern learner \( \mathcal{L}_P \) uses inductive program synthesis [32] for synthesizing patterns that describe a given set \( S \) of strings using a user-specified set of atoms \( \mathcal{A} \). For convenience of end users, we automatically enrich their specified set of atoms by allowing: (1) all Const atoms, and (2) fixed-width variants of all Class atoms specified by them, for all widths. \( \mathcal{L}_P \) is instantiated with these enriched atoms derived from \( \mathcal{A} \), which we denote as \( \langle \mathcal{A} \rangle \):

\[
\langle \mathcal{A} \rangle = \mathcal{A} \cup \{\text{Const} \mid s \in S\} \cup \{\text{Class} \mid \text{Class}_z \in \mathcal{A} \land z \in \mathbb{N}\}
\]

(1)

Although \( \langle \mathcal{A} \rangle \) is unbounded (since \( S \) is unbounded), as we explain later, our synthesis procedure only explores a small fraction of \( \langle \mathcal{A} \rangle \) which are compatible with a given dataset \( S \).

We build on a state-of-the-art inductive program synthesis library PROSE [4], which implements the FlashMeta [40] framework. Our synthesis relies on deductive reasoning – reducing a synthesis problem over an expression, to smaller synthesis problems over its subexpressions. PROSE provides a convenient framework with highly efficient algorithms and data-structures for building such program synthesizers.

An inductive program synthesis task is defined by: (1) a domain-specific language (DSL) for target programs which in our case is \( \mathcal{L}_P \), (2) a spec [40, §3.2] that defines a set of constraints over the output of the desired program. The spec given to \( \mathcal{L}_P \) (denoted as \( \varphi \)), simply requires the desired pattern \( P \) to describe all strings i.e. \( \forall s \in S: P[s] = \text{True} \).

We formally write such a spec \( \varphi \) as:

\[
\varphi \overset{\text{def}}{=} \bigwedge_{s \in S} [s \rightarrow \text{True}]
\]

Deductive reasoning allows us to reduce the spec \( \varphi \) over a pattern \( P \) to specs over its arguments. The specs are reduced recursively till terminal symbols in \( \mathcal{L}_P \) (string \( s \), or atom \( \alpha \)). Then, starting from the values of terminal symbols which satisfy their spec, we collect the subexpressions and combine them to synthesize bigger expressions. We refer the reader to [40] for more details on the synthesis process.

The reduction logic (called witness functions [40, §5.2]) for specs is domain-specific, and depends on the semantics of the DSL operators. Specifically for \( \mathcal{L}_P \), we need to define the logic for reducing the spec \( \varphi \) over the two kinds of patterns: Empty, and \( P[\text{SuffixAfter}(s, \alpha)] \).

For \( P[\text{Empty}(s, \alpha)] \) to satisfy \( \varphi \), i.e., describe all strings \( s \in S \), each string \( s \) must indeed be \( \epsilon \). No further reduction is needed since \( s \) is a terminal. We simply check, \( \forall s \in S: s = \epsilon \).

\( P[\text{Empty}(s, \alpha)] \) fails to satisfy \( \varphi \) if \( \varphi \) contains non-empty strings.

\[
P[\text{SuffixAfter}(s, \alpha)] \overset{\text{def}}{=} \bigwedge_{s \in S} [s[\alpha(s);] \rightarrow \text{True}]
\]

(2)

Instead of enumerating \( \varphi_{\alpha} \) for all allowed atoms, we consider only those atoms \( \alpha \) that succeed in matching some prefix of all strings in \( S \). This is the key to computing the intersection of patterns over individual strings in an incremental fashion. Such atoms are said to be compatible with the given dataset \( S \).

**Definition 5.3.** Compatible Atoms: Over a set \( U \) of allowed atoms, we say a set of atoms \( \mathcal{A} \subseteq U \) is compatible with a given dataset \( S \), denoted as \( \mathcal{A} \propto S \), if every atom in \( \mathcal{A} \) successfully matches some prefix of every string \( s \in S \), i.e.

\[
\mathcal{A} \propto S \iff \forall \alpha \in \mathcal{A}: \forall s \in S: \alpha(s) > 0
\]

We call a compatible set \( \mathcal{A} \) of atoms is maximal under \( \propto \), denoted as \( \mathcal{A} = \max_{\propto}^S \{ \mathcal{A} \} \). We refer the reader to [43] for a formal definition.

Example 9. Consider the following Cardinal postal codes: \( S = \{ \text{V6E3V6, V6C2S6, V6X1X5, V6X3S4} \} \). Over the set \( \langle \mathcal{A} \rangle \) of 18 atoms, we can implement the set \( \Lambda = \max_{\propto}^S \{ \mathcal{A} \} \) using equation (1) on \( \mathcal{A} = \text{default atoms listed in Table 1} \), the maximal set of compatible atoms \( \max_{\propto}^S \{ \mathcal{A} \} \) contains 18 atoms such as "V6", "V", "Upper", "Upper+", "Alpha+", "AlphaSpace", "AlphaDigit" etc.

**GetMaxCompatibleAtoms**, outlined in Figure 11, accepts a set of strings \( S \), a set of atoms \( A \) specified by the user, and computes the set \( \Lambda = \max_{\propto}^S \{ \mathcal{A} \} \), where \( \mathcal{A} \) denotes the enriched set of atoms based on \( \mathcal{A} \) given by equation (1). We start with \( \Lambda = \mathcal{A} \) (line 1), and gradually remove atoms that are not compatible with \( S \), i.e., fail to match at least one string \( s \in S \) (line 5). For enriching \( A \) with fixed-width Class tokens, we maintain a hashtable \( C \) that maps a Class token to its width (line 8). If the width of a Class atom is not constant over all \( s \in S \), we remove it from \( C \) (line 9). We finally add to \( \Lambda \), the fixed-width variants of all Class atoms in \( C \), generated by calling \( \text{RestrictWidth}(\cdot) \) (line 10). For enriching with Const atoms, we compute the longest common prefix \( L \) across all \( s \in S \) (line 11), and add every prefix of it to \( \Lambda \), as a compatible Const token (line 12).

In essence, a spec \( \varphi \) for \( P[\text{SuffixAfter}(s, \alpha)] \) is reduced to a set of \( \max_{\propto}^S \{ \mathcal{A} \} \) new specs, each representing a distinct synthesis problem for \( P: \{ \varphi_{\alpha} \mid \alpha \in \max_{\propto}^S \{ \mathcal{A} \} \} \), where \( \varphi_{\alpha} \) is defined as equation (2), and \( \mathcal{A} \) denotes the enriched set of atoms derived from \( \mathcal{A} \) by equation (1). Each synthesis problem is solved similarly as the original one. PROSE handles the recursive propagation of appropriate specs to subexpressions, and combines the generated subexpressions to patterns that satisfy the original spec \( \varphi \).

We conclude with a comment on the soundness and completeness of our pattern learner \( \mathcal{L}_P \).
Definition 5.4. Soundness and U-Completeness: We say that a learner \( \mathcal{L}_P \) for \( \mathcal{L}_P \) patterns is sound, if for a given dataset \( S \) every learned pattern \( P \) satisfies \( \forall s \in S : P[s] = \text{true} \). We say that \( \mathcal{L}_P \) instantiated with a set \( U \) of atoms is U-complete, if for every dataset \( S \), it learns all patterns \( P \in \mathcal{L}_P \) over \( U \) atoms, which satisfy \( \forall s \in S : P[s] = \text{true} \).

For our synthesis procedure, soundness is guaranteed since we only consider compatible atoms. Completeness follows from the fact that we always consider their maximal set over \( U \) atoms. \( \mathcal{L}_P \) is \( \langle A \rangle \)-complete for any user-specified set \( A \) of atoms. Therefore, once the set of patterns \( \mathcal{L}_P(S) \) has been learned for a \( S \), a variety of cost functions may be employed to select the most suitable pattern for \( S \) amongst all possible patterns over \( \langle A \rangle \), without recomputing \( \mathcal{L}_P(S) \).

5.3 Cost of Patterns in \( \mathcal{L}_P \)

Our cost function \( C_P \) produces a real number, given a pattern \( P \in \mathcal{L}_P \) and a dataset \( S \), based on the structure of \( P \) and its behaviour over \( S \). Empty is assigned a cost of 0 regardless of the dataset, since any dataset with which \( C_P \) is consistent, only contain \( \epsilon \). We define the cost \( C_P(P, S) \) for a pattern \( P = \alpha_1 \cdot \cdots \cdot \alpha_n \) with respect to dataset \( S \) as:

\[
C_P(P, S) = \sum_{i=1}^{n} C_P(\alpha_i) \cdot D(i, P, S)
\]

Each atom \( \alpha \) in \( \mathcal{L}_P \) has a statically assigned cost \( C_P(\alpha) \in (0, C_{max}) \) based on a priori bias for the atom. The static costs for the default atoms in \( \text{FlashProfile} \) were empirically decided. We define \( C_{max} = C_P(\text{Any}^+) \) to be the maximum cost across all atoms. For a pattern \( P = \alpha_1 \cdot \cdots \cdot \alpha_n \), our cost function \( C_P \) sums these static costs after applying a dynamically determined weight \( D(i, P, S) \in (0, 1) \), based on how well each token \( \alpha_i \) generalizes over \( S \). 

\[
D(i, P, S) = \frac{1}{|S|} \sum_{s_j \in S} l_{i,j}
\]

where \( l_{i,j} = (l_{j,1}, \ldots , l_{j,n}) \) denotes the lengths of successive prefix matches over a string \( s_j \in S \). Since a token match never fails over \( S \) for any synthesized pattern \( P \), \( l_{i,j} > 0 \) and \( D(i, P, S) > 0 \) for all tokens \( \alpha_i \in P \).

Example 10. Consider \( S = \{ \text{Male}, \text{Female} \} \), that are matched by \( P_1 = \text{Upper} \cdot \text{Lower}^+ \) and \( P_2 = \text{Upper} \cdot \text{HexDigit} \cdot \text{Lower}^+ \). The static costs for the relevant atoms are:

\[
\{\text{Upper} \rightarrow 8.2, \text{HexDigit} \rightarrow 26.3, \text{Lower}^+ \rightarrow 9.1\}
\]

The costs for both patterns shown above, are computed as:

\[
C_P(P_1, S) = \frac{8.2}{2}(\frac{1}{4} + \frac{1}{6}) + \frac{26.3}{3}(\frac{1}{2} + \frac{1}{3}) + \frac{9.1}{3}(\frac{2}{4} + \frac{2}{5}) = 8.9
\]

\[
C_P(P_2, S) = \frac{6.2}{2}(\frac{1}{4} + \frac{1}{8}) + \frac{28.3}{3}(\frac{1}{2} + \frac{1}{3}) + \frac{9.1}{3}(\frac{2}{4} + \frac{2}{5}) = 12.5
\]

\( P_1 \) is chosen as best pattern, since \( C_P(P_1, S) < C_P(P_2, S) \).

Note that although \text{HexDigit} is more specific compared to \text{Upper} and \text{Lower} - \text{HexDigit} contains 16 characters as opposed to 26, it has a higher static cost. This is an example of a priori bias against \text{HexDigit} to avoid strings like “face” being described as \text{HexDigit} instead of \text{Lower}.

However, its cost is much lower compared to \( C_P(\text{AlphaDigit}) = 639.6 \), making it the preferred pattern for strings such as “408d”. Static costs are baked into the atoms and must be provided by domain experts when introducing new atoms.

\( C_P \) balances the trade-off between specificity vs generality - more specific atoms receive a smaller dynamic weight (which leads to a smaller overall cost), whereas the sum of the costs over many overly specific atoms may exceed the cost of a single more general atom. In § 6.2, we evaluate the quality of profiles learned by \text{FlashProfile} and show that they are natural - neither too specific, nor overly general.

6. EVALUATION

In this section, we present an experimental evaluation of FlashProfile, focusing on the following key questions:

- §6.1 How well does our syntactic similarity measure perform over real world entities?
- §6.2 How accurate are the profiles over real datasets, and what is the effect of sampling and approximations?
- §6.3 What is the overall performance of FlashProfile, and how does it depend on the various parameters?
- §6.4 Are the generated profiles natural and useful? How do they compare to those from existing tools?

Implementation and Experimental Setup: We have implemented FlashProfile as a cross-platform C# library built using Microsoft PROSE [4]. All experiments were performed on an 8-core Intel i7 3.6GHz machine with 32GB RAM running 64-bit Ubuntu 16.10 with .NET Core 1.0.1.

6.1 Syntactic Similarity

We evaluate the applicability of our dissimilarity measure from Definition 4.1, over real-life entities. From 25 clean real datasets\(^{11}\) ranging over names, dates, postal codes, phone numbers etc. in different formats, we randomly pick two datasets, and select a random string from each. We picked 240400 such pairs of strings. A good similarity measure is expected to be able to identify when the two strings are drawn from the same dataset by assigning them a lower dissimilarity value, compared to two strings selected from different datasets. For example, the pair \( \{\text{Albert Einstein}, \text{Isaac Newton}\} \) should have a lower dissimilarity value than \( \{\text{Albert Einstein}, 03/20/1998\} \).

For evaluation, we use the standard precision-recall [35] (PR) measure. Precision in our context is the fraction of pairs that truly belong to the same dataset, out of all pairs that are labeled to be “similar” by the predictor. Recall is the fraction of pairs retrieved by the predictor, out of all pairs truly drawn from same datasets. By varying the threshold for labelling a pair as “similar”, we generate a PR curve and measure the area under the curve (called AUC). A good similarity measure should exhibit high precision and high recall, and therefore have a high AUC. Figure 12 show a comparison of our method against two

\(^{10}\)Available at: https://github.com/SaswatPadhi/ProfilingExperiments

\(^{11}\)Available at: https://github.com/SaswatPadhi/ProfilingExperiments
baselines: (1) a character-based similarity measure (JarW), and (2) a machine-learned predictor (RF) using several intuitive syntactic features. We explain them below.

We observed that character-based measures [17] show poor AUC, and are not indicative of syntactic similarity. A popular data-wrangling tool OpenRefine [3] allows clustering of string data using Levenshtein distance [30]. However, this measure exhibits a negligible AUC over our benchmarks.

Although the Jaro-Winkler distance [49] indicated as JarW in Figure 12(a) shows a better AUC, it is quite low compared to both our, and machine-learned predictors.

Our second baseline is a standard random forest [11] model RF using the syntactic features listed in Figure 12(b) such as, difference in length, number of digits etc. We train RF over 160,267 pairs with $(\frac{1}{2})^2 = 0.16\%$ pairs drawn from same datasets. We observe from Figure 12(a) that the accuracy of RF is quite susceptible to changes in the distribution of the training data. RF2 and RF3 were trained with 0.64% and 1.28% pairs from same datasets, respectively. While RF3 performs marginally better than our predictor, RF3 performs worse. In contrast, our technique does not require any training, and hence even if training and test distributions are significantly different, our method still produces an accurate similarity measure.

6.2 Profiling Accuracy

We show FlashProfile’s accuracy along two dimensions:

• **Partitions:** Our sampling and approximation techniques preserve accuracy of partitioning the datasets.

• **Descriptions:** Profiles generated using $\mathcal{D}_P$ and $\mathcal{C}_P$ are natural – neither too general nor too specific.

**Partitioning:** From 25 clean datasets, we randomly pick $n \in \{2, 8\}$ datasets, and select a group of 256 random strings from each. We combine them, and invoke FlashProfile to partition them into $n$ clusters. We measure the precision of clustering using the symmetric uncertainty [50], which is a measure of normalized mutual information (NMI). An NMI of 1 indicates the resulting partitioning to be identical to the original groupings of strings, and an NMI of 0 indicates that the final partitioning is unrelated to the original groupings.

Recall that in each iteration, we profile $|M|$ strings, and sample pairwise dissimilarities of only $|P| = |\theta M|$ strings w.r.t. $S$. For this experiment, we set $M = n$. With different values of $\mu$ and $\theta$, we show the mean NMI of the partitionings over 10 tests for each value of $n$, in Figure 13(a).

The NMI improves with increasing $\theta$, since we sample more dissimilarities, resulting in better approximations. However, the NMI drops with increasing $\mu$, since more pairwise dissimilarities are approximated. We observe that the median NMI is significantly higher than the mean, indicating a small number of cases where FlashProfile made poor approximations. The dashed line indicates the median NMI with $\mu = 4.0$. We observe a median NMI of 0.96 (mean 0.88) for $(\mu = 4.0, \theta = 1.25)$, which is FlashProfile’s default configuration for our experiments (indicated by the circled point in Figure 13(a)).

**Descriptions:** We evaluate the suitability of the automatically suggested profiles, by measuring their precision and recall. A natural profile should not be overly specific – it should generalize well over the dataset (true positives), but not beyond it (false positives).

We consider 56 datasets ignoring datasets with duplicate formats. For each dataset, we profile a randomly selected subset containing 10% of its strings, and measure: (1) the fraction of the remaining dataset described by it, and (2) the fraction of an equal number of strings from other datasets, matched by it. Figure 13(b) summarizes our results. The lighter and darker shades indicate the fraction of true positives and false positives respectively. The white area at the top indicates the fraction of false negatives – the fraction of the remaining dataset that is not described by the profile. We record an overall precision of 97.8%, a recall of 93.4%. The dashed line indicates a mean true positive rate of 93.2%, and the dotted line shows a mean false positive rate of 2.3%.

6.3 Performance

We evaluate FlashProfile’s performance over various $(\mu, \theta)$-configurations considered during the partitioning-accuracy evaluation. We show the performance-accuracy trade off in Figure 14(b). We plot the mean speed up of various configurations over $(\mu = 1.0, \theta = 1.0)$, against the mean NMI of partitioning. Our default configuration $(\mu = 4.0, \theta = 1.25)$ achieves a mean speed up of 2.3×. We also show the profiling times in Figure 14(a). The dotted lines indicate profiling time without pattern sampling, for different values of the string-sampling factor $\mu$. The dashed line shows the median profiling time for various values of $\theta$ with $\mu = 4$.

As one would expect, the profiling time increases with $\theta$, since FlashProfile samples more patterns making more calls to $\mathcal{D}_P$. The dependence of profiling time on $\mu$ however, is more interesting. Notice that for $\mu = 1$, the profiling time is higher than any other configuration.
with pattern sampling enabled (solid lines). This is due to the fact that FlashProfile learns very specific profiles with \( \mu = 1 \) over a very small sample of strings, and does not describe much of the remaining data. This results in many Sample → Profile → Filter iterations over the entire dataset. The performance improves with \( \mu \) till \( \mu = 4.0 \), and then starts deteriorating as we sample more pair wise dissimilarities.

Finally, we evaluate FlashProfile’s performance on end-to-end real-life profiling tasks on 74 datasets collected from various online sources, that have a mixture of clean and dirty datasets. Over 142 tasks – 74 for automatic profiling, and 68 for refinement, we observe a median profiling time of 0.7s. With our default configuration, 77% of the requests and 68 for refinement, we observe a median profiling time of 0.7s. With our default configuration, 77% of the requests and 68 for refinement, we observe a median profiling time of 0.7s.

![Figure 15: Profiling time for real-life datasets.](image)

7. APPLICATIONS IN PBE SYSTEMS

In this section, we briefly discuss applications of our data profiling technique to improve PBE systems. Such systems aim to synthesize a desired program from a small number of input-output examples [18, 21, 45]. For instance, given an example “Albert Einstein” → “A.E.”, the goal is to learn a program that extracts the initials for a given name. That is, given “Alan Turing”, we want the synthesized program to output “A.T.”. Though several PBE systems have been proposed recently, a major criticism for these systems has been the lack of usability and confidence in them [28, 37].

Examples are an inherently under-constrained form of specs for the desired program, and a large number of programs (up to \( 10^{20} \)) may be consistent with them [41]. Two major challenges to learning the desired program are: (1) obtaining a representative set of examples that convey the desired behavior, and (2) ranking the consistent programs to select the ones natural to end users.

**Significant Inputs:** A user of the PBE system cannot be expected to provide the representative examples. In fact, typically, users provide outputs for only the first few inputs. However, if all these examples are very similar, the system may not learn a program that generalizes over other inputs. In §2, we discuss such a case for Flash Fill [18], a popular PBE system for string transformations. Instead, we propose to request the user to provide outputs for significant inputs – the most dissimilar input w.r.t. those previously provided.

First, we generate a syntactic profile \( P \) for the input, and use the OrderP Arts function, listed in Figure 16(a), to order the input partitions, based on mutual dissimilarity. Starting with the tightest partition (line 1) i.e. the one with the least-cost pattern describing it, we iteratively append
the partition that is most dissimilar (requires the highest-cost pattern to describe) with prior partitions in $\rho$ (line 3). We request the user to provide an output for a randomly selected input from each partition in order $\langle S_1, \ldots, S_{|P|} \rangle$. Since Flash Fill is interactive, the user can inspect the output for each input, and skip if it is already correct. Finally, we restart from $S_1$ after one cycle through all partitions.

We measure the efficacy of our interaction model, over 163 Flash Fill benchmarks that require $> 1$ examples to learn the desired program. Figure 16(b) compares the number of examples required originally, to that using significant inputs. Seven benchmarks that timed-out have been omitted. Over the remaining 156 benchmarks, we observe that Flash Fill, (1) never requires a second example from the same partition, for 131 benchmarks, and (2) uses the smallest possible set of examples over the given inputs, for 140 benchmarks.

Thus, (1) validates our hypothesis that syntactic partitions indeed identify representative inputs, and (2) further indicates that ordering partitions using our dissimilarity measure, allows for a highly effective interaction model.

Note that, the significant inputs scenario is similar to active learning, which is well-studied in machine-learning literature [23]. Active learning also seeks to find data points to be annotated so that the learned predictor is most accurate. However, typical active-learning methods require hundreds of annotated examples. In contrast, PBE systems typically deal with very few annotated examples [37].

8. RELATED WORK

Data Profiling: There has been a line of work on profiling various aspects of a column of data; see [38, 6] for recent surveys. Traditional profiling techniques target simple statistical properties [36, 12, 13, 24, 3, 5].

To our knowledge, no existing technique supports refinement of syntactic profiles learned over an extensible language. We present a novel dissimilarity measure which is the key to learning refinable profiles over user-specified patterns. While Potter’s Wheel [43] does not learn a compile profile, it learns the most frequent data pattern using user-defined domains. SSDT [2] learns rich regular expressions but is neither extensible not comprehensive. A dedicated profiling tool Ataccama One [1] generates comprehensive profiles over a very small set of base patterns. OpenRefine [3] does not learn syntactic profiles, but it allows clustering of strings using character-based similarity measures [17]. In §6 we show that they do not capture syntactic similarity.

Application-Specific Structure Learning: There has been prior work on learning specific structural properties aimed at aiding data wrangling applications, such as data transformations [43, 45], information extraction [31], and reformatting or normalization [44, 27]. These approaches make specific assumptions regarding the target application, which do not necessarily hold when learning general purpose comprehensive profiles for data understanding. We show in §7 that profiles learned by FlashProfile may aid PBE based applications, such as Flash Fill [19] for data transformation.

A recent work leverages profiling based on hierarchical clustering, for tagging sensors used in building automation [10]. However, they use a fixed set of features relevant to their domain, and do not qualify clusters with patterns.

Grammar Induction: Syntactic profiling is also related to the problem of learning regular expressions, or more generally a grammar [14] from a given set of examples. Most techniques in this line of work such as L-Star [7] and RPNI [39], assume availability of both positive and negative examples, or a membership oracle. Furthermore, these techniques are either too slow or do not generalize well [9].

Finally, the LearnPADS [16, 51] tool generates a syntactic description and a suite of tools for processing semi-structured data. However it does not support refinement.

Program Synthesis: Our implementation, FlashProfile, uses an inductive program synthesis framework, PROSE [4] (also called FlashMeta [40]). Inductive synthesis, specifically programming-by-examples [19, 21] (PBE) has been the focus of several recent works on automating data-driven tasks, such as string transformations [18, 46, 45], text extraction [29], and spreadsheet data manipulation [20, 22]. However, unlike these applications, data profiling does not solicit any examples from the user. We demonstrate a novel application of a supervised synthesis technique to solve an unsupervised learning problem — our clustering technique drives the synthesizer by creating examples as necessary.

9. CONCLUSION

With increasing volume and variety of data, we require better profiling techniques to enable end users to easily understand, and analyse their data. Existing techniques target simple data-types, mostly numeric data, or only generate partial profiles for string data, such as frequent patterns. In this paper, we present a technique for learning comprehensive syntactic descriptions of string datasets, which also allows for interactive refinement. We implement this technique as FlashProfile, and present extensive evaluation on its accuracy and performance. We show that profiles generated by FlashProfile are useful both for manual data analysis and in existing PBE systems.

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