The Problem Solving Genome: Analyzing Sequential Patterns of Student Work with Parameterized Exercises

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ABSTRACT

Parameterized exercises have recently emerged as an important tool for online assessment and learning. The ability to generate multiple versions of the same exercise with different parameters helps to support learning-by-doing and decreases cheating during assessment. At the same time, our experience with using parameterized exercises for Java programming reveals suboptimal use of this technology as demonstrated by repeated successful and failed attempts to solve the same problem. In this paper we present the results of our work on modeling and examining patterns of student behavior with parameterized exercises using Problem Solving Genome, a compact encapsulation of individual behavior patterns. We started with micro-patterns (genes) that describe small chunks of repetitive behavior and constructed individual genomes as frequency profiles that shows the dominance of each gene in individual behavior. The exploration of student genomes revealed that individual genome is very stable, distinguishing students from their peers and changing very little with the growth of knowledge over the course. Using the genome, we were able to analyze student behavior on the group level and identify genes associated with good and bad learning performance.

Categories and Subject Descriptors
Information systems [Information Systems Applications]; Data mining

Keywords
sequential pattern mining, parameterized exercises

1. INTRODUCTION

Parameterized exercises have recently emerged as an important tool for online assessment and learning. A parameterized exercise is essentially an exercise template that is instantiated at runtime with randomly generated parameters. As a result, a single template is able to produce a large number of similar, but distinct questions. While parameterized questions are considerably harder to implement than traditional “static” questions, the benefits offered by this technology make this additional investment worthwhile. During assessment, a reasonably small number of question templates can be used to produce online individualized assessments for large classes minimizing cheating problems [12]. In a self-assessment context, the same question can be used again and again with different parameters, allowing every student to achieve understanding and mastery. The above mentioned properties of parameterized exercises made them very attractive for the large-scale online learning context. At the same time, parameterized exercises as a learning technology have its own problems. Our experience with personalized exercises for SQL [17] and Java [7] in the self-assessment context demonstrated that the important ability to try the same question again and again is not always beneficial, especially for students who are not good in managing their learning. The analysis of a large number of student logs revealed some considerable number of unproductive repetitions. We observed many cases where students kept solving the same exercise correctly again and again with different parameters, well passed the point when it could offer any educational benefit. While it might increase self-confidence, students’ time and effort might be spent better by advancing to more challenging questions. We also observe cases where students persisted in failing to solve the same, too difficult exercise, instead of focusing on filling the apparent knowledge gap or switching to simpler exercises.

The work presented in this paper was motivated by our belief that the educational value of parameterized exercises could be increased by a personalized guidance mechanism that can predict non-productive behavior and intercept it by recommending a more efficient learning path. Main challenge with predicting unproductive behavior is to examine the stability of behavior patterns in the problem solving process. If the patterns, such as specific unproductive sequences, appear at random, there is a slim chance to predict and prevent them. If, on the contrary, specific patterns are associated with certain features of the student (such as knowledge and individual traits), exercise complexity, or the learning process stage, there is a good chance to learn the association rules and use it for prediction. In this paper we performed an extended study of problem solving patterns in the con-
text of parameterized exercises. We explored the connection between these patterns and the components of the learning process mentioned above. Our study produced a rather unusual result. While it was more plausible to expect that the patterns are related to the current level of student knowledge, our analyses revealed that the patterns are related to student problem solving tendency. More exactly, we discovered that every student has a specific combination of micro-patterns, a kind of problem solving genome. We observed that this genome is relatively stable, distinguishing every student from his or her peers, it changes very little with the growth of the student knowledge over the course. We also discovered that genomes are not randomly distributed, and instead, students with similar genomes form cohorts that perform relatively similarly in the problem solving process. We believe that our discovery of problem solving genome is a very important step towards our goal of predicting and preventing unproductive behavior. Indeed, the stability of patterns on the personal level makes the task of pattern prediction feasible while the presence of cohorts opens the way to detect student problem-solving genome early in the learning process. In this paper we present our approach of detecting student problem-solving genome and report our exploration of the genome on the level of individual students and cohorts.

The rest of the paper is structured as follows. Next section briefly reviews several areas of related work. Section 3 describes the dataset used in the study. Section 4 presented the method for building the Problem Solving Genome. In Section 5 we explore the Genome stability and it’s relation with performance groups and the complexity of the exercises. Section 6 summarizes the contribution and discusses future work.

2. RELATED WORK

2.1 Parameterized Questions and Exercises

Recent studies in educational technology have demonstrated promising results by leveraging computer and Web abilities to deliver parameterized exercises worldwide, which has become one of the focusing topics in Web-enhanced education. One of the most influential system, CAPA [9], was evaluated in a number of careful studies [8, 9], providing clear evidence that individualized exercises can significantly reduce cheating while improving student understanding and exam performance. The CAPA technology has been later integrated into popular LON-CAPA platform [12] and its functionality defined the assessment architecture of eDX. Due to the complexity of parameterized assessment, the majority of work on parameterized questions and exercises was done in physics and other math-related domains where a correct answer to a parameterized question can be calculated by a formula. There are, however, examples of using this technology in other domains. In particular, our team focused on parameterized exercises for teaching programming. We developed and explored QuizPACK platform for C-programming [3] and a similar QuizJET platform for Java programming [7]. Problem solving repetition behaviors has been studied by psychologists in different ways, providing evidence that repetition behaviors have roots in cognitive, metacognitive and motivational aspects and explaining why some students quit and some persist when facing challenging problems [14]. Schunk [16] shows the positive correlation between persistency in repeating and self-efficacy (believe on self capabilities/skills to solve a problem). The attribution theory [19] describes how students that attribute performance outcomes (successes, failures) to effort tend to work harder than students who attribute them to ability. Grounded in the literature in educational psychology, we conjecture that patterns on problem solving repetition may be explained by individual learners’ motivational traits that are part of learners’ personality [15]. These theories provide insights into analyzing to which extent these behaviors are stable on students.

2.2 Sequential Pattern Mining in Educational Context

Mining sequential patterns of students actions has recently gained attention in educational data mining field. Using activity data collected from groups of students working with interactive tabletops, Martínez et al [13], mined and clustered frequent patterns to compare distinct behaviors between low and high achievement groups. The differential sequence mining method, introduced by Kinnebrew and Biswas [11] has been successfully used to differentiate behavioral patterns among groups of students (such as low and high performance students). The method uses SPAM [1] to find common patterns in the sequences of the whole dataset, and then applies statistical tests to reveal differences in the frequencies of the discovered patterns among different groups. The same authors have applied this technique in data collected from the system Betty’s Brain to discovered patterns that can distinguish self-regulated behaviors in successful and non-successful students [2], and to analyze the evolution of reading behaviors in high and low performance students during productive and non-productive phases of work [10]. Herold, Zundel and Stahovich [4] have used the differential sequence mining on sequences of actions on handwritten tasks and proposed a model to predict performance on the course based on pattern features. Our work extends this prior work by utilizing and aggregating the mined sequence patterns to construct student activity profiles. Such profiles enable us to evaluate the statistical differences at the student, exercise, and group levels.

3. SYSTEM AND DATASET

We collected answers of students who worked with QuizJET [7] parameterized Java exercises in the context of an introductory object-oriented programming class at the School of Information Sciences in the University of Pittsburgh. The students accessed the exercises through Progressor+ interface [6]. The system was provided for self-study and its use was not mandatory. Each QuizJET exercise was generated from a template by substituting a parameter variable with a randomly generated value. Exercises generated using the same template were equal from semantics point of view. To answer the exercise the student had to mentally execute a fragment of Java code to determine a value of a specific variable or the content printed on a console. When the user answers, the system evaluates the correctness, reports to the student whether the answer was correct or wrong, shows the correct response, and invites the student to “try again”. Next time, the exercise will be generated with other values and the correct answer will be different. In this way, the student can try the same exercise many times, leaving a trace
of successes and failures. In total, Progressor+ provided access to 103 different parameterized exercises organized in 19 topics (Variables, Objects, Arrays, etc.). Exercises are also labeled in terms of complexity as easy, medium and hard. There are 41 easy exercises, 41 medium exercises and 19 hard exercises. The dataset includes three semesters of student data (Spring 2012, Fall 2012 and Spring 2013). Overall, the dataset recorded 6489 incorrect and 14726 correct attempts. Easy exercises were attempted 10620 times, medium complexity exercises were attempted 7876, and hard exercises were attempted 2719 times. Once started to work with an exercise the students might attempt it just once or try it several times in a sequence. The dataset includes 4212 single attempts (no repetition) and 4758 sequences with more than 1 attempt. Among these there are 2717 with more than 2 attempts, 1583 with more than 3 attempts, and 1016 sequences with more than 4 attempts.

4. BUILDING THE PROBLEM SOLVING GENOME

The key idea of our “genome” approach is to build a compact characteristics of student problem-solving behavior on the level of micro-patterns. To build a genome we started with finding proper micro-patterns (genes) and then built a genome of a student as a vector representing the frequencies of different micro-pattern occurrences in the student problem-solving logs. An overview of the genome-building process is shown in Figure 1. To build the genes, we started by label students’ attempts using time and correctness (Figure 1(a), Section 4.1). We then apply sequential pattern mining to extract sequential micro-patterns Figure 1 (b), Section 4.2. Most frequent micro-patterns were selected as genes and used as a basis for the Problem Solving Genome, which is a vector of gene frequencies (Figure 1(c), Section 4.3). This section presents the genome-building process in details while the next sections report our exploration of the Genome.

4.1 Attempts labeling

We use both time and correctness of each attempt to label it for further use in sequential pattern mining analysis. In this way, each action will convey more information than using correctness only. As shown in the Figure 2, distribution of times for first attempts are different from other (non-first) attempts. This is reasonable if we consider that the user needs extra time the first time to read and understand the exercise. Additionally, time distribution is different for different exercises, as in general, complex exercises need longer times. Thus, for labeling the time factor, we used time information of historical records in our system to compute median times for each exercise for both first and other attempts. Then, we labeled the attempt as short or long depending on the time being lower or greater than the median of the distribution for the specific exercise. Combining correctness and time, we finally label the attempts using the letters ‘s’ (lowercase s) for a short success, ‘S’ (uppercase S) for a long Success, ‘f’ for a short failure, ‘F’ for a long Failure.

The labeled attempts are organized in sequences by pairs student-question within a session in the system. Each sequence s_u,e represents the sequential attempts of user u in the exercise e within a session. If the user attempted the same exercise in different sessions, there will be more than one sequence s_u,e. Additionally, we mark starting and ending points on sequences using ‘_’ (underscore). For example, a sequence _fSs_s means start with a short failure, make a long success and then finish with a short success.

4.2 Sequential pattern mining

To discover frequent patterns, we use PexSPAM algorithm [5], which extends the fast SPAM algorithm [1] with gap and regular expression constraints. Given a sequence database \( D = s_1, s_2, ..., s_n \), the support of a pattern \( \alpha \) is the number of sequences of \( D \) which contains \( \alpha \) as a subsequence at least once. If the support of \( \alpha \) is bigger than a threshold, then \( \alpha \) is considered a frequent pattern. Support measure does not inform for multiple occurrences of the pattern within a sequence. In this work, we set a small minimum support in 1% because even when a pattern occurs in overall few sequences, it can still make a difference when looking at the aggregation of pattern occurrences by pattern. Additionally, since we are interested in looking at patterns of 2 or more sequential attempts, we set the gap in 0 and considered only sequences with more than 1 attempt. After running the mining algorithm, we discover 102 common patterns occurring at least in 1% of the sequences. These common micro-patterns of student behavior play the role of genes in our approach. The top 20 genes and the corresponding support can be seen in Table 1.
4.3 The problem solving genome: characterizing students with pattern vectors

Using the 102 gene patterns discovered by the sequential pattern mining, we build individual frequency vectors that show how frequently each gene appears in student problem solving behavior. Since this vector captures in a compact form the specifics of student problem solving behavior, we call it student Problem Solving Genome. Note that frequency-based approach allows building individual genome using any subset of gene sequences, for example, all sequences in the term, the first half of sequences of the student activity on the term, a random subset of sequences, etc.

Since a pattern might occur more than once in a sequence, and more than one pattern may occur in a sequence, the frequency vectors are not summing to 1. Thus, we normalize the vectors for further analysis.

5. EXPLORING THE PROBLEM-SOLVING GENOME

5.1 Problem Solving Genome stability

The first step of problem-solving genome exploration is assessing its stability. To what extent the name “genome” that we assigned to the micro-pattern frequency vector is justified? Is it just a random mix of pattern which could be different for different time slots or, like a real genome, it is a stable characteristic of a user that distinguishes him or her from the peers? A good approach to check genome stability is to randomly split sequences of user activity patterns into two equal sets and build the genome vector from each of two halves. If the genome is stable, then two random halves of the split genome should be significantly closer to each other than to half-genomes of other users. In contrast, if genome halves are no closer to each other than to half-genome vectors of other users, we can’t consider genomes as stable user characteristics. To assess the stability hypothesis we built two half-genomes for each user by randomly splitting his or her observed sequences in half and compiling gene frequency vectors for each half. We then calculate pairwise distances between all half-genomes.

To compute distances, we use Jensen-Shannon (JS) divergence as it is a symmetric version of Kullback-Leibler divergence and has been widely used for computing distance between frequency distributions. Additionally, we filter out all student with less than 60 sequences, limiting differences due to extreme differences on amount of activity. There are 32 students with at least 60 sequences. In this analysis we use paired samples t-test on the difference between the self and other distances. Normality assumption is met. Results are shown in Table 2 first row (a). Students self-distances are significantly smaller ($M = .2370, SE = .0169$) than distances to other students ($M = .4815, SE = .0141$), $t = -15.224, p < .001$, Cohen’s $d = 2.693$.

While similarity of random half-genomes is a very strong argument in favor of genome stability, the random split has one weak aspect: since each of the random halves represents student micro patterns over the whole duration of the course, it is still possible that student genome gradually changes over the course duration from one pattern frequency to another. To assess temporal stability of genome we need to use temporal split, i.e., to compare half-genomes built from the temporally first half (early) and second half (late) of student sequences. Results on Table 2 second row (b) confirm the temporal stability hypothesis: distances between halves of the same genome ($M = .3211, SE = .0214$) are significantly smaller than between-student distances ($M = .4997, SE = .0164$), $t = -6815, p < .001$, Cohen’s $d = 1.205$. This result is very important, it confirms that individual problem solving genome is stable, it characterizes each user as individual and doesn’t change with the growth of his or her knowledge or course progression. In other words, the frequencies of micro-pattern appearances is a true “genome” that uniquely characterizes every user while sufficiently distinguishing them from others.

5.2 Effect of complexity

While we discovered that the knowledge level and course stage doesn’t affect the genome, it is still possible that behavior patterns are affected by exercise complexity. To understand how the complexity level of the exercises impact on the pattern frequencies, we analyze distances between the genome of the exercises (i.e. pattern frequency vector for each exercise). Having the exercises’ genome and the predefined classification in easy, medium and hard, we select pairs of exercises within and between complexity levels and compute distances using Jensen-Shannon divergence. We filter out all questions with less than 20 sequences and perform comparisons between extremes groups, i.e. easy and hard complexity levels to extreme the differences. Normality and homogeneity of the variance on pair distances are not met on all levels, thus non-parametric test is applied. Results of the Kruskal-Wallis test shows significant differences between distances within and between levels, $\chi^2(2, N = 1596) = 160.359, p < .001$. Mann-Whitney test is performed to test differences among the levels. Distances within easy exercises (mean rank = 626.16) are significantly smaller than distances between easy and hard exercises (mean rank = 909.77), $z = -12.564, p < .001$. Similarly, the distances within hard exercises (mean rank = 277.20) are significantly smaller than distances between easy and hard exercises (mean rank = 383.13), $z = -4.733, p < .001$. These results show a clear dependency of the pattern behaviors with the complexity level of the questions. This is reasonable given that hard questions, which need more time, are expected to discourage repetitions.

The impact of exercise difficulty on the behavior patterns leaves open an interesting opportunity that genome is as much impacted by the unique exercise difficulty profile or every user as by their individual differences. To exclude this option, we re-examine the analysis on Section 5.1 now considering randomly split genome built only from activity on easy exercises, to control for differences of students amount

Table 3: Mean and standard error of distances within and between easy and hard exercises.

<table>
<thead>
<tr>
<th></th>
<th>Mean</th>
<th>SE</th>
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<tbody>
<tr>
<td>within easy</td>
<td>.3311</td>
<td>.0031</td>
</tr>
<tr>
<td>within hard</td>
<td>.3478</td>
<td>.0085</td>
</tr>
<tr>
<td>between easy-hard</td>
<td>.4145</td>
<td>.0050</td>
</tr>
</tbody>
</table>
of activity on different complexity exercises. We perform this analysis with 39 students having at least 20 sequences in easy questions. Results shown in last row (c) in Table 2 confirm the stability of patterns: students are more similar to themselves (self distance $M = .3736$, $SE = .0214$) than to others (distances $M = .6065$, $SE = .0128$), $t = -10.352$, $p < .001$, Cohen’s $d = 1.6569$, even within the exercises of the same complexity.

### 5.3 Patterns of Success within student groups

Since one of the goals of this paper is using behavior analysis to identify and prevent inefficient patterns, it would be valuable to use the genome to identify which patterns make groups of students more or less successful in the learning process. The easiest approach to do it is to split students into performance-related groups and find unique genome aspects in this group. This simple approach, however, might not work since for students with very different genomes, different behavior patterns might be related to success. In this case, to find connection between patterns and performance, we should group students into groups with similar behavior and contrast most and least successful students within each group. In this section we perform both kinds of the analysis.

#### 5.3.1 Behavior Patterns for Predefined Performance Groups

Predefined Performance Groups (PPG) are defined based on pre and posttest scores that we collected. The pre and posttest were highly similar among different semesters (small variation on questions) and the scores were further normalized as (score) / (max score) (having that min score is 0). Additionally, using the normalized pre and posttest scores, we compute a normalized learning gain score as (normalized post score) - (normalized pre score). For each of the pretest, posttest, and learning gain measures, students are classified in three groups using the percentiles 33.3 and 66.7: low, medium, and high. For example, a student with pretest lower or equal than the percentile 33.3 in the pretest score distribution is classified as low pretest student. Summarizing, we have 3 PPG (low, medium, high) for each performance measure (pretest, posttest, and learnig gain).

Table 2 shows the number of students in each predefined performance group (PPG).

Table 4: Number of students in each predefined performance group (PPG).

<table>
<thead>
<tr>
<th>Group</th>
<th>Pretest (total=67)</th>
<th>Posttest (total=65)</th>
<th>Learning gain (total=65)</th>
</tr>
</thead>
<tbody>
<tr>
<td>low</td>
<td>24</td>
<td>22</td>
<td>22</td>
</tr>
<tr>
<td>medium</td>
<td>16</td>
<td>19</td>
<td>20</td>
</tr>
<tr>
<td>high</td>
<td>27</td>
<td>24</td>
<td>23</td>
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</tbody>
</table>

We collected 97 pretest and 93 posttest results in the 3 semester. We further filter out the students with few usage of the system, setting a threshold of minimum 20 sequences and minimum 2 sessions. Additionally, we exclude one student that present in the first 6 sequences a very unusual number of repetitions, and we consider this student as a clear outlier. At the end our data consists in 67 students having pretest, and 65 of them having both pre and posttest.

Does students with similar performances have similar patterns for solving parameterized exercises? Is this similarity, between the students of the same predefined performance group, more than the similarity we can find between the students from different groups? For this analysis we contrast the genome built using all the term activity (all problem solving sequences) of the students classified in the performance groups described before. We sample 50% of all possible pairs of students within and between PPGs and compute the distances (Jensen-Shannon divergence) of all within and between group pairs. Then, we compare the average of distances within and between groups to see if students inside each group are more similar to each other than to students in other groups. Normality and homogeneity of variance is not met for all groups, thus we use Krustal-Wallis non-parametric mean rank test and Mann-Whitney test for single comparisons. We constrained the analysis to PPGs low and high to see extreme differences.

Results are shown in Table 5. Mann-Whitney comparison is reported only where significant differences among groups were found (pretest). For pretest groups, distances within the low group (mean rank = 222.70) are significantly smaller than distances between low and high groups (mean rank = 258.21), $z = -2.537$, $p = .011$. This suggests that student with no previous experience tend to behave differently than students with stronger background. There is no significant difference between high and low-high distances, though, meaning that high group behave more heterogeneously than low group. For posttest and learning gain groups there are no significant differences on distances within and between groups. This results are intriguing, as we will expect to find clear differences among performance groups. Since we could not find those differences, we could hypothesize that specific behavior patterns can’t be easily characterized as universally helpful or harmful for student performance, instead, the impact of each micro-pattern on student behavior might depend on the whole profile of micro-patters, i.e., the genome. This, to find connections between genome and performance, we need to start from the opposite side: cluster the students based on the genome, characterize the clusters in terms of the distinguishable patterns, and find helpful and harmful patterns within each class. We describe these analyses in the following sub sections.

#### 5.3.2 Clustering students by their genome

We use the genome as a feature vector and cluster students using spectral clustering technique [18] as it gives a better separation of the students. We choose two clusters (K=2) as we observe that two clusters give the largest eigen-gap,
sustaining there are two intrinsic groups in the data. Figure 3 shows the top 30 frequent patterns in both of the clusters. Each point represents the average frequency of seeing a particular pattern in the cluster. Error bars are included to indicate significance. We order the patterns in x-axis by the differences between clusters 2 and 1. As we can see in this figure, some of the patterns, such as $\text{FS}$, $\text{FS}$, $\text{ss}$, $\text{SS}$, etc., occur with significant frequency difference in the two clusters and some other patterns, such as $\text{fS}$, $\text{fS}$, $\text{Fs}$, etc., do not show significant differences. If we look more closely, the sequences that start with failure are mostly related to the students in cluster 2 and the sequences that start with success are mostly related to the students in cluster 1. Also, we can see that the students in cluster 1 tend to repeat their successful attempts more and more frequently (e.g., the $\text{ssss}$ pattern). In other words, even when they get the right answer to the question, they will insist on confirming knowing the question by repeating it again and again. Unlike students in cluster 1, the students in cluster 2 are much less prone to this “confirmation” behavior. Instead, they are more prone to stop working with an exercise early, frequently right after figuring out the first right answer to the question, even if they have struggled for the correct answer in their previous attempts (e.g., $\text{FS}$, $\text{FS}$, and $\text{FS}$ patterns). Thus, using the student genome, we can identify two major types of student behaviors in solving parameterized exercises. Based on these observations above, we call the first cluster of the students the confirmer and the second cluster the non-confirmers.

### 5.3.3 Performance differences among clusters

Once two clusters of students that are similar in their overall behavior are identified, we can re-examine the connection between student success in the course and the aspects of their genomes on the cluster level. To find out the cluster-level relationship between different behaviors and learning outcomes, we study pattern by pattern differences between different PPGs within each cluster and describe the patterns that distinguish them. Both of the clusters have students from all PPGs. As a result, we cannot say that the student’s genome has a direct impact on the performance of the student. Both confirmer and non-confirmers can have high or low performance. To look at the clusters deeply and to see if there are any differences in the patterns, within each cluster, that can drive students’ performance, we repeat the first analysis within each cluster looking at the learning gain. For each of the clusters, we look at the patterns and the difference between their average frequencies for the students with low and high learning gain. The result is shown in Figure 4. The upper diagram shows the students in cluster 1 (the confirmer) and the lower diagram shows the students in cluster 2 (the non-confirmers). The red line with round markers show the pattern frequencies for low learning gain students and the blue line with the triangle marker is representative of high learning gain students.

If we look at the patterns in cluster 1 (the confirmer), we can see that there are some patterns that show significant difference between the low and high learning gain students. All of these patterns start with a failure: $\text{FS}$ and $\text{Ff}$ have long failures in the beginning of the patterns and $\text{FF}$, $\text{fS}$, and $\text{ff}$ have short failures at the beginning of the patterns. Among these patterns, only $\text{FS}$ is practiced more by the high learning gain students. This indicates that, among the confirmer students, the ones that put a good amount of effort to answer a question right, after a long failure and stop repeating the same question learn more. The low learning gain group shows more frequent use of the $\text{ff}$, $\text{fS}$, and $\text{ff}$ patterns. The common element of all of these patterns is short failure ($\text{f}$). If we look at Figure 4 for confirmer, we can see that all of the patterns that include a short failure, are practiced more by the low gain students. This can indicate

<table>
<thead>
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<th>Table 5: Statistical tests on differences on distances between pairs of students within low, within high, and between low and high PPGs.</th>
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<tr>
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<tr>
<td>---------------------------------------------------------------</td>
</tr>
<tr>
<td>Pretest</td>
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<tr>
<td>Posttest</td>
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<td>L. Gain</td>
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</table>

**Figure 3:** Top 30 patterns and their frequencies in each cluster. Patterns are ordered by the difference on frequencies between cluster 2 (non-confirmers) and 1 (confirmers).
that the low gain confirmers students do not spend enough
time and thought on the questions that they do not know
the answer of.

The non-confirmers show more pattern differences between
the low and high learning gainers. We can see that the high
learning gain group follow the patterns of _FF, FS, fFS,
SS, SS, SS, and Ss more frequently. This means that the
high learning gain, non-confirmers students tend to continue
trying a non-parameterized exercise and spending time on it
after they failed in it or it took them a long time to get to the
correct answer for that exercise. In this sense, these students
are closer to the confirmers group of students (cluster 1) but
only at the times that they are not sure if they have learnt
the solution to an exercise. On the other hand, the low
learning gain group tend to develop the fs, fs, and ff
patterns in their sequences. The first two indicates that they
give up practicing the exercise after having a short success
that comes after a short failure. Also, they tend to repeat
short failures on the same exercise more often.

Comparing beneficial and harmful patterns for the two clus-
ters, we can make an interesting observation that the in-
creased use of several beneficial patterns for each cluster
make students more familiar to the opposite cluster. For ex-
ample, while confirmers have generally low tendency to stop
after first hard success _FS_ successful confirmers demon-
strate this pattern much more frequently. On the other side,
while non-confirmers generally tend to stop after first hard
success, successful non-confirmers have higher tendency to
continue after hard success as shown by significantly in-
creased frequencies of such patterns as SS, SS, and Ss
In other words, while the two clusters are considerably dif-
ferent by their behavior overall, the “centrist” students that
are closer to the opposite cluster tend to be more successful,
while the extreme behavior that distinguishes the cluster is
frequently related to less successful performance.

Another interesting observation here is that having repeated
successes in the same exercise does not add to the learning
gain of the students. We can see that none of the patterns
having more than one short success make any significant
differences between the low and high learning gain students.
The above analysis shows the specific patterns that can ex-
plain the differences between high and low learning gain stu-
dents in each of the confirmers and non-confirmers clusters.
In both of the clusters, short failures are more associated
with low learning gain students. For the non-confirmers
cluster group, the students, who acted similar to the confirmers
group in cases of having a hard time getting to the right
answer, have higher learning gain. Also, repeating the short
success did not add to the learning gain of students. These
results are promising for the further guidance of the stu-
dents in the correct use of the system to increase their per-
f ormance. Based on a students’ pattern cluster, we can en-
courage them to follow the sequences that are associated
with high learning gain for their cluster (such as encourag-
ing them to think longer on questions) and discourage them
from following the patterns that have no effect or negative
effect (e.g. stopping the student from repeating short suc-
cesses).

6. CONCLUSIONS AND FUTURE WORK

In this paper we explored patterns of student repetitive work
with parameterized exercise for Java programming domain.
The goal of this work was to understand the connections
between micro- and macro-level behavior patterns and fac-
tors that might be responsible for this behavior such as ex-
ercise difficulty, student personality, level of knowledge, or
position in the course. In turn, we hoped that this under-
standing could help us predict how a specific student would
work with a specific exercise and prevent inefficient behav-
ior such as repetitive successful attempts to solve an exercise
when the exercise become too easy to contribute to student
knowledge growth. To explore the impact of students’ per-
sonal features on their work with programming exercises,
we build the student problem solving genome, a compact
representation that encapsulates the specifics of individual
behavior patterns. To build the genome, we started with
micro-patterns (genes) that describe small chunks of repet-
itive behavior in relation to the correctness and duration of
each attempt. We then constructed a genome as a frequency
profile that shows the dominance of each gene in the student
behavior.

Using the genome approach we analyzed the stability of be-
havior patterns for students and groups and explored their
connection with student success in the course. The most
interesting finding was stability of the genome on individual level. As our analysis showed, the genome characterizes a user as a person rather than her level of knowledge as might be expected in an educational system. It uniquely identifies a user among other users and doesn’t change with the considerable growth of student knowledge over the course duration. While the problem complexity does affect the behavior patterns as well, we demonstrated that the genome is defined by some inherent characteristics of the user rather than a difficulty profile of the problems they solve.

To find connection between problem-solving genome and student performance, we examined genomes for various groups of students. Since a direct attempt to associate genome with performance-related groups (a typical way group students in educational context) was not successful, we started from the opposite side and formed student groups on the basis of their genome (i.e., behavior) similarity. As it appears, all students could be most reliably split into just two cohorts that are differ considerably by their behavior. After that split, we were able to contrast successful and less successful learners by their behavior and identify “beneficial” and “harmful” genes for each cohort. In particular, it was interesting to observe that the behavior of successful learners in each cohort was somewhat closer to the behavior of the opposite cohort.

In the future work we would like to proceed to our ultimate goal of recognition and prediction of inefficient behavior. The discovery of a stable genome provides a good ground for developing a recognition engine and the presence of behavior cohorts indicates that some good guidance (encouraging “beneficial” micro-patterns and discouraging “harmful” ones) could be provided even in the early stage of student work when it might be harder to build a reliable genomic profile. We also believe that the “genome” approach provides a new way for exploration of student problem solving behavior and plan to explore to the stability of “genome” and the presence of behavior cohorts in other domains with parameterized problem solving.

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8. REFERENCES