# Algorithms for Computing Approximate Repetitions in Musical Sequences

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**Abstract.** Here we introduce two new notions of approximate matching with application in computer assisted music analysis. We present algorithms for each notion of approximation: for approximate string matching and for computing approximate squares.

**Keywords**: String algorithms, approximate string matching, dynamic programming, computer-assisted music analysis.

### 1 Introduction

This paper focuses on a set of string pattern-matching problems that arise in musical analysis, and especially in musical information retrieval. A musical score can be viewed as a string: at a very rudimentary level, the alphabet could simply be the set of notes in the chromatic or diatonic notation, or the set of intervals that appear between notes (e.g. pitch may be represented as MIDI numbers and

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pitch intervals as number of semitones). Approximate repetitions in one or more musical works play a crucial role in discovering similarities between different musical entities and may be used for establishing 'characteristic signatures' (see [6]). Such algorithms can be particularly useful for melody identification and musical retrieval.

Both exact and approximate matching techniques have been used for a variety of musical applications (see overviews in McGettrick [24]; Crawford et al [6]; Rolland et al [29]; Cambouropoulos et al [4]). The specific problem studied in this paper is pattern-matching for numeric strings where a certain tolerance is allowed during the matching procedure. This type of pattern-matching has been considered necessary for various musical applications and has been used by some researchers (see, for instance, Cope [5]). A number of efficient algorithms will be presented in this paper that tackle various aspects of this problem.

Most computer-aided musical applications adopt an absolute numeric pitch representation (most commonly MIDI pitch and pitch intervals in semitones; duration is also encoded in a numeric form). The absolute pitch encoding, however, may be insufficient for applications in tonal music as it disregards tonal qualities of pitches and pitch-intervals (e.g. a tonal transposition from a major to a minor key results in a different encoding of the musical passage and thus exact matching cannot detect the similarity between the two passages). One way to account for similarity between closely related but non-identical musical strings is to use what will be referred to as  $\delta$ -approximate matching (and  $\gamma$ -approximate matching). In  $\delta$ -approximate matching, equal-length patterns consisting of integers match if each corresponding integer differs by not more than  $\delta$ - e.g. a C-major  $\{60, 64, 65, 67\}$  and a C-minor  $\{60, 63, 65, 67\}$  sequence can be matched if a tolerance  $\delta = 1$  is allowed in the matching process ( $\gamma$ -approximate matching is described in the next section). Two simple musical examples that illustrate the usefulness of the proposed pattern-matching techniques are presented in Appendices I and II.

Exact repetitions have been studied extensively. The repetitions can be either concatenated with the original substring or they may overlap or they may not. Algorithms for finding non-overlapping repetitions in a given string can be found in [?,8,15,22,19,27] and algorithms for computing overlapping repetitions can be found in [3,13,14,26]. A natural extension of the repetitions problem is to allow the presence of errors; that is, the identification of substrings that are duplicated to within a certain tolerance k (usually edit distance or Hamming distance). Moreover, the repeated substring may be subject to other constraints: it may be required to be of at least a certain length, and certain positions in it may be required to be invariant.

Furthermore, efficient algorithms for computing the approximate repetitions are also directly applicable to molecular biology (see [11, 18, 25]) and in particular in DNA sequencing by hybridization ([28]), reconstruction of DNA sequences from known DNA fragments (see [17, 30]), in human organ and bone marrow transplantation as well as the determination of evolutionary trees among distinct species ([17]).

Another type of repetition that is used in computer assisted music analysis is that of finding evolutionary chains: given a string t ("the text") and a pattern p (the "motif"), find whether there exists a sequence  $u_1 = p, u_2, \ldots, u_l$  occurring in the text t such that  $u_{i+1}$  occurs to the right of  $u_i$  in t and  $u_i$  and  $u_{i+1}$  are "similar" (i.e. the differ by a certain number of symbols). In [9] and [7] algorithms for overlapping and non-overlapping evolutionary chains were presented and several variants of the problem were studied: computing the longest chain, computing the chain with the least number of errors.

The paper is organised as follows. In the next section we present some basic definitions for strings and background notions for approximate matching. In Section 3 we present an algorithm for  $\delta$ -approximate (the first notion of approximation) pattern matching. In section 4 we present an algorithm for  $\delta$ ,  $\gamma$ -approximate (the second notion of approximation) pattern matching. In section 5 we present algorithms for computing all  $\delta$  and  $\{\delta,\gamma\}$ - approximate squares in a given text. Finally in Section 6 we present our conclusions and open problems.

# 2 Background and basic string definitions

A string is a sequence of zero or more symbols from an alphabet  $\Sigma$ ; the string with zero symbols is denoted by  $\epsilon$ . The set of all strings over the alphabet  $\Sigma$  is denoted by  $\Sigma^*$ . A string x of length n is represented by  $x_1 \ldots x_n$ , where  $x_i \in \Sigma$  for  $1 \leq i \leq n$ . A string w is a substring of x if x = uwv for  $u, v \in \Sigma^*$ ; we equivalently say that the string w occurs at position |u| + 1 of the string x. The position |u| + 1 is said to be the starting position of w in x and the position |w| + |u| the end position of u in x. A string w is a prefix of x if x = wu for  $u \in \Sigma^*$ . Similarly, w is a suffix of x if x = uw for  $u \in \Sigma^*$ .

The string xy is a concatenation of two strings x and y. The concatenations of k copies of x is denoted by  $x^k$ . For two strings  $x=x_1\ldots x_n$  and  $y=y_1\ldots y_m$  such that  $x_{n-i+1}\ldots x_n=y_1\ldots y_i$  for some  $i\geq 1$ , the string  $x_1\ldots x_ny_{i+1}\ldots y_m$  is a superposition of x and y. We say that x and y overlap.

Let x be a string of length n. A prefix  $x_1 ldots x_p$ ,  $1 \le p < n$ , of x is a period of x if  $x_i = x_{i+p}$  for all  $1 \le i \le n-p$ . The period of a string x is the shortest period of x. A string y is a border of x if y is a prefix and a suffix of x.

Let  $\Sigma$  be an alphabet of integers. We say that two symbols a,b are  $\delta$ -approximate, denoted  $a=_{\delta}b$  if and only if

$$|a-b| \leq \delta$$

where  $\delta$  is a given integer. We say that two strings x, y are  $\delta$ -approximate, denoted  $x \stackrel{\delta}{=} y$  if and only if

$$|x| = |y|$$
, and  $x_i = y_i, \forall i \in \{1..|x|\}$  (2.1)

for a given integer  $\delta$ .

We say that two strings x, y are  $\gamma$ -approximate, denoted  $x \stackrel{\gamma}{=} y$  if and only if

$$|x| = |y|$$
, and  $\sum_{i=1}^{|x|} |x_i - y_i| < \gamma$  (2.2)

for a given integer  $\gamma$ . Furthermore, we say that two strings x,y are  $\{\gamma,\delta\}$ -approximate, denoted  $x\stackrel{\gamma,\delta}{=} y$ , if and only if x and y satisfy conditions (2.1) and (2.2).

# 3 $\delta$ -Approximate Pattern Matching

The problem of  $\delta$ -approximate pattern matching is formally defined as follows: given a string  $t = t_1 \dots t_n$  and a pattern  $p = p_1 \dots p_m$  compute all positions j of t such that

$$p \stackrel{\delta}{=} t[j..j + m - 1]$$

The algorithm is based on the O(1)-time computation of the "Delta states"  $DState_j, j \in \{1..n\}$  by using bit operations under the assumption that  $m \leq w$ , where w is the number of bits in a machine word. The basic steps of the algorithm are as follows:

- 1. First we compute the "Delta table" DT: we set  $DT(\alpha) = r$ , where  $\alpha$  denotes a symbol occurring in t and  $r = r_1 \dots r_m$  is a binary word with  $r_i$  equal to 1 if  $|\alpha p_i| \leq \delta$ , otherwise  $r_i$  is equal to 0 for  $i \in \{1..m\}$ .
- 2. Let LeftShift be a bit-wise operation that shifts the bits of a binary word by one position to the left. We define

$$DState_j := (LeftShift(DState_{j-1}) \text{ OR } 1) \text{ AND } DT[t_j]$$
 (3.1)

for j=1...n and  $DState_0=0$ ; hence this procedure is called "Shift-And". Once we have computed the DT table, we can use it to compute the  $DState_j$  for j=1...n, using the recursive formula (3.1).

3. We say that there is a  $\delta$ -approximate match (or simply  $\delta$ -match) at position j-m+1 if and only if the m-th bit of  $DState_j$  is 1 or equivalently if and only if  $DState_j$ , is greater or equal to  $2^{m-1}$  when it is viewed as a decimal integer.

Example. For  $\Sigma = \{1, \ldots, 9\}$  let us consider  $p=3,4,6,2,\ t=3,4,6,2,8,2,4,5,7,1$  and  $\delta = 1$ . In the preprocessing table,  $DT(\alpha)$  denotes the positions where  $|\alpha - p_i| \le \delta$ . For example, DT[3] = 1011 because  $|3 - p_i| \le 1$  for i=1,2,4.

The table below evaluates  $DState_i$  using the relation (3.1). For example,

 $DState_4 = (LeftShift(DState_3) \text{ OR 1}) \text{ AND } DT[t_4]$ 

= (LeftShift(0100) OR 1) AND DT[2]

= (1000 OR 1) AND 1001

= 1001 AND 1001

= 1001

i	$p_i$	DT[1]	DT[2]	DT[3]	DT[4]	DT[5]	DT[6]	DT[7]	DT[8]	DT[9]
4	2	1	1	1	0	0	0	0	0	0
3	6	0	0	0	0	1	1	1	0	0
2	4	0	0	1	1	1	0	0	0	0
1	3	0	1	1	1	0	0	0	0	0

**Table 1.** The table DT for pattern p = 2, 6, 4, 3 and alphabet  $\Sigma = \{1, \dots, 9\}$ .

j	1	2	3	4	5	6	7	8	9	10
$t_{j}$	3	4	6	2	8	2	4	5	7	1
$LeftShift(DState_{j-1}) \text{ OR } 1$	0001	0011	0111	1001	0011	0001	0011	0111	11 01	1001
$DT[t_j]$	1011	0011	0100	1001	0000	1001	0011	0110	0100	1000
$DState_j$	0001	0011	0100	1001	0000	0001	0011	0110	0100	1000
$[DState_j]_{10}$	1	3	4	9	0	1	3	6	4	8

**Table 2.** Computing the *Dstates* and finding the  $\delta$ -approximate matches.

which implies that there is a match starting at position 1 of t, since the 4-th bit of  $DState_4$  is 1.

A  $\delta$ -approximate match occurs at position j-m+1 of t if  $[DState_j]_{10} \geq 2^{m-1}$ , where  $[DState_j]_{10}$  denotes the  $DState_j$  as a decimal integer. Therefore, there is one match ending at position 4 of t ( $\{3,4,6,2\}$ ) and another one at position 10 of t ( $\{4,5,7,1\}$ ) since  $\{DState_4, DState_{10}\} \geq 2^3$ .

#### 3.1 Pseudo-code

Fig. 1 gives a complete specification of the algorithm. In the line 3 we have the preprocessing phase which compute the DT table. In line 6 we use the recursive formula to compute the DStates. Finally, in line 7 we apply the matching criteria to see whether there is a  $\delta$ -approximate match or not.

#### 3.2 Running time

Assuming that the pattern length is no longer than the memory word size of the machine (thus O(1) size), the time complexity of the preprocessing phase is O(n) (since we need to evaluate DT only for the symbols that occur in t) and the time complexity of the searching phase in O(n). Figure 2 shows the timing <sup>1</sup> for different text sizes.

# 4 $\{\delta, \gamma\}$ -Approximate Pattern Matching

The problem of  $\{\delta, \gamma\}$ -approximate pattern matching is formally defined as follows: given a string  $t = t_1 \dots t_n$  and a pattern  $p = p_1 \dots p_m$  compute all positions

<sup>&</sup>lt;sup>1</sup> Using a SUN Ultra Enterprise 300MHz running Solaris Unix.

```
1. procedure SHIFT-AND(p,\,t,\,\delta) { n=|t|,\,m=|p| }  
2. begin  
3. DT_i[\alpha] = \begin{cases} 1 & \text{if } |\alpha-p_i| \leq \delta \\ 0 & \text{otherwise} \end{cases} \forall i \in \{1..m\}, \; \forall \alpha \in \Sigma  
4. DState_0 \leftarrow 0  
5. for j \leftarrow 1 to n do  
6. DState_j \leftarrow (LeftShift(DState_{j-1}) \text{ OR } 1 \text{ ) } \text{ AND } DT[t_j]  
7. if DState_j \geq 2^{m-1} then write j-m+1  
8. od  
9. end
```

Fig. 1. The Shift-And Procedure.

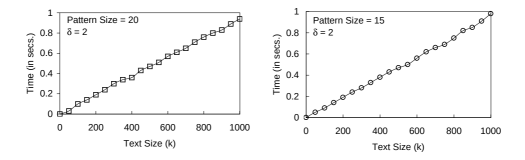


Fig. 2. Timing curves for the Shift-And Procedure.

j of t such that

$$p \stackrel{\delta,\gamma}{=} t[j..j + m - 1]$$

In order to solve this problem we first make use of the Shift-And algorithm to find the  $\delta$ -approximate matches of the pattern p in t. Once we find a  $\delta$ -approximate match we want to know whether it is also a  $\gamma$ -approximate match. To do so, we seek to compute successive "Delta States"  $DState_j$  and "Gamma States"  $GStates_j$  in O(1) time using bit operations under the assumption that  $m \leq w$  where w is the number of bits in a machine word. The main step of the algorithm are as follows:

- 1. We need to compute the "Delta Table" DT as we did before and the "Gamma Table" GT table; we set  $GT(\alpha) = r$ , where  $\alpha$  denotes a symbol in the alphabet and  $r = r_1, \ldots, r_m$  is a word with  $r_i$  equal to  $|\alpha p_i|$  if  $|\alpha p_i| \leq \delta$ , otherwise  $r_i$  is equal to 0 for  $i \in \{1..m\}$ . Each  $r_i$ ,  $i \in \{1..m\}$  is stored as a binary number of d bits where  $d = \lceil log(\delta \times m) \rceil$ .
- 2. Let LeftShift be a bit-wise operation that shifts the bits of a binary word one position to the left and RightShift shifts the bits of a binary word d positions to the right. Once we have computed the DT and GT tables, we

can use them to compute the  $DState_j$  and  $GState_j$  for j=1...n, using the recursive formulas

$$DState_{i} = (LeftShift(DState_{i-1}) \ OR \ 1) \ AND \ DT[t_{i}]$$
 (4.1)

and

$$GState_{j} = RightShift(GState_{j-1}, d) + GT[t_{j}]$$
(4.2)

We also need to define the seeds  $DState_0=0$  and  $GState_0=0$ . We call this procedure "Shift-Plus" because we use the "shift" and "plus" operators to compute each new state.

3. We say that there is a match  $(\{\delta, \gamma\}$ -approximate match) at position j-m+1 if and only if the m-th bit of  $DState_j$  is 1 and the m-th block of d bits taken as an integer is  $\leq \gamma$ .

Example. For our example let  $\Sigma = \{1, \ldots, 9\}$ , the pattern p = 3, 4, 6, 2, the text t = 3, 4, 6, 2, 8, 2, 4, 5, 7, 1,  $\delta = 1$  and  $\gamma = 3$ . We will use blocks of size three (d = 3) to store the  $|\alpha - p_i|$  values where  $|\alpha - p_i| \leq \delta$ . For example,  $GT[3] = 000\ 100\ 000\ 100$  because  $|3 - p_i| \leq 1$  for i=1,2,4 and the differences are 0,1,1 respectively. (see left hand table of table 3).

i	$p_i$	1	2	3	4	5	6	7	8	9
		0	1	0	1	0	0	0	0	0
1	3	0	0	0	0	0	0	0	0	0
		0	0	0	0	0	0	0	0	0
		0	0	1	0	1	0	0	0	0
2	4	0	0	0	0	0	0	0	0	0
		0	0	0	0	0	0	0	0	0
		0	0	0	0	1	0	1	0	0
3	6	0	0	0	0	0	0	0	0	0
		0	0	0	0	0	0	0	0	0
		1	0	1	0	0	0	0	0	0
4	2	0	0	0	0	0	0	0	0	0
		0	0	0	0	0	0	0	0	0

j	1	2	3	4	5	6	7	8	9	10
$t_{j}$	3	4	6	2	8	2	4	5	7	1
	0	1	0	1	0	1	1	0	0	0
3	0	0	0	0	0	0	0	0	0	0
	0	0	0	0	0	0	0	0	0	0
	1	0	1	0	1	0	1	0	0	0
4	0	0	0	0	0	0	0	1	0	0
	0	0	0	0	0	0	0	0	0	0
	0	1	0	1	0	1	0	0	1	0
6	0	0	0	0	0	0	0	1	1	0
	0	0	0	0	0	0	0	0	0	0
	1	0	1	0	1	0	1	0	0	0
2	0	0	0	0	0	0	0	0	1	0
	0	0	0	0	0	0	0	0	0	1

**Table 3.** The left hand side table is the "Gamma Table" GT and the right hand side table is the table for finding  $\{\gamma, \delta\}$ -approximate matches.

The right hand table above shows the computation of the DStates and the GStates using (4.2). For example,

```
GState_9 = RightShift(\ 000010010000,3\ ) + 000000100000 \\ = 000000010010 + 000000100000 = 000000110010
```

We already know that there are two  $\delta$ -approximate matches ending at positions 4 and 10 of t. Now we can use the last three bits of  $GState_4$  and  $GState_{10}$  to find out the values of  $\gamma$ , which are 0 and 4 respectively (see right hand table of Fig. 3).

#### 4.1 Pseudo-code

Fig. 3 below gives a complete description of the algorithm. In the lines 3 and 4 are the preprocessing phase which compute the DT table and GT table respectively. In lines 8 and 9 we compute the next DState and GState respectively. Finally, in line 10 we apply the matching criteria to see whether there is a match or not.

```
procedure SHIFT-PLUS(p,\,t,\,\delta,\,\gamma) { n=|t|,\,\,m=|p| }
1.
2.
               DT_{i}[\alpha] = \begin{cases} 1 & \text{if } |\alpha - p_{i}| \leq \delta \\ 0 & \text{otherwise} \end{cases} \quad \forall i \in \{1..m\}, \forall \alpha \in \Sigma 
GT_{di-d...di-1}[\alpha] = \begin{cases} |\alpha - p_{i}| & \text{if } DT_{i}[\alpha] = 1 \\ 0 & \text{otherwise} \end{cases} \quad \forall i \in \{1..m\}, \forall \alpha \in \Sigma 
3.
4.
                DState_0 \leftarrow 0
5.
                GState_0 \leftarrow 0
6.
                for j \leftarrow 1 to n do
7.
                         DState_i \leftarrow (LeftShift(DState_{i-1}) \text{ OR 1}) \text{ AND } DT[t_i]
8.
                         GState_j \leftarrow RightShift(GState_{j-1}, d) + GT[t_j]
9.
                         if DState_j \ge 2^{m-1} AND GState_{dm-d...dm-1} \le \gamma then write j-m+1
10.
11.
                od
12.
          end
```

Fig. 3. The SHIFT-PLUS Algorithm.

# 4.2 Running time

Assuming that  $\delta \times m \leq 2^d-1$  the time complexity of the preprocessing phase is  $O(\delta \times m + |\Sigma|)$  and the time complexity of the searching phase in O(n), thus independent from the alphabet size and the pattern length. Figure 4 shows the timing for different text sizes.

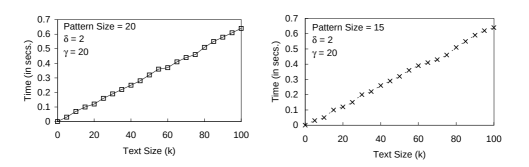


Fig. 4. Timing curves for the Shift-Plus Algorithm.

# 5 Computing Approximate Squares

The problem of computing all  $\delta$ -approximate squares is formally defined as follows: given a string  $t = t_1 \dots t_n$  and an integer  $\delta$ , compute all positions j of t for which  $\exists u \in \Sigma^m$  such that

$$t[j..j+m] \stackrel{\delta}{=} u$$
 and  $t[j+m+1..j+2m] \stackrel{\delta}{=} u$ 

The string u is said to be the root of the square.

The problem of computing all  $\{\delta, \gamma\}$ -approximate squares is formally defined as follows: given a string  $t = t_1 \dots t_n$  and an integer  $\delta$ , compute all positions j of t for which  $\exists u \in \Sigma^m$  such that

$$t[j..j+m] \stackrel{\delta,\gamma}{=} u$$
 and  $t[j+m+1..j+2m] \stackrel{\delta,\gamma}{=} u$ 

where u is said to be the root of the square.

When we look for a square we will run into two possibilities: One when the root occurs in the square and other when the root does not occurs necessarily in the square.

# 5.1 Consider an approximate square such that the root occurs in the square

The diagonal diag(i) corresponds to the pair of positions (j, j + i) and therefore to the candidates for squares of length 2i. There exists an approximate square of length 2i at position j if there exists a run of values not greater than  $\delta$  of length at least i on the diagonal diag(i) starting at position j.

For example, consider diag(2) (see table 4) and  $\delta = 1$ . We are trying to locate runs of length at least 2 containing only values not greater than  $\delta = 1$ . We obtain:

Position	Square	$\operatorname{Roots}$				
14	(2,3,1,4)	(2,3) or $(1,4)$				

For this example we only have a  $\delta$ -approximate square starting at position 14 which root can be either (2,3) or (1,4). Note that the roots certainly occur in the square.

# 5.2 Consider an approximate square such that the root does not occur necessarily in the string

We say that there exists an approximate square of length 2i at position j if there exists a run of values not greater than  $2\delta$  of length at least i on the diagonal diag(i) starting at position j. In other words, we are using  $2\delta$  instead of  $\delta$ .

For example, consider diag(2) (see table of Fig. 4) and  $\delta = 1$ . We are trying to locate runs of length at least 2 containing only values not greater than  $2\delta = 2$ . We obtain:

$\begin{array}{ c c c c c c c c c c c c c c c c c c c$							1			5	_		_	_	10				1.5			19		
2       -3       -5 <th c<="" td=""><td></td><td></td><td></td><td></td><td><math>\boldsymbol{x}</math></td><td>=</td><td>2 -</td><td>-3 -</td><td><math>5  ext{ } 4</math></td><td>-1</td><td>-7</td><td>1</td><td>-5</td><td>-5</td><td>3</td><td>-3</td><td>1</td><td>1 2</td><td>3</td><td>1</td><td>4 5</td><td>5 7</td><td>_</td></th>	<td></td> <td></td> <td></td> <td></td> <td><math>\boldsymbol{x}</math></td> <td>=</td> <td>2 -</td> <td>-3 -</td> <td><math>5  ext{ } 4</math></td> <td>-1</td> <td>-7</td> <td>1</td> <td>-5</td> <td>-5</td> <td>3</td> <td>-3</td> <td>1</td> <td>1 2</td> <td>3</td> <td>1</td> <td>4 5</td> <td>5 7</td> <td>_</td>					$\boldsymbol{x}$	=	2 -	-3 -	$5  ext{ } 4$	-1	-7	1	-5	-5	3	-3	1	1 2	3	1	4 5	5 7	_
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$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	15	3		-	-	-	-	-	-	•	•	-	-	-	-	-	-	0	2	1	2	4	$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	
$  \cdot \cdot \cdot   \cdot   \cdot   \cdot   \cdot   \cdot   \cdot   \cdot   \cdot$	16	1	1	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	0	3	4	6	$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	
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	18	5	1	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	0	2	$\setminus \text{diag}(2)$	
19 7 $-1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1$	19	7	1	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	0	$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	
$\wedge$ diag $(0)$								•					•					•		-	_	-	$\setminus \operatorname{diag}(0)$	

Table 4. Table for computing approximate squares.

Position	Square	Roots
5	(-1,-7,1,-5)	(0,-6)
9	(-5,3,-3,1)	(-4,2)
12	(1,1,2,3)	(1,2) or $(2,2)$
13	(1,2,3,1)	(2,1) or $(2,2)$
14	(2,3,1,4)	(1,3), (1,4), (2,3)  or  (2,4)

Furthermore, consider diag(3) and  $\delta=1$ . We are trying to locate runs of length at least 3 containing only values not greater than  $2\delta=2$ . We obtain:

Position	Square	Roots
1	(2,-3,-5,4,-1,-7)	(3,-2,-6)
6	(-7,1,-5,-5,3,-3)	(-6,2,-4)
12	(1,1,2,3,1,4)	(2,0,3), (2,1,3)  or  (2,2,3)
13	(1,2,3,1,4,5)	(0,3,4), (1,3,4)  or  (2,3,4)

In those cases where we want to consider a  $\{\delta, \gamma\}$ -approximate square we just check each  $\delta$ -approximate match to see if it is also a  $\{\delta, \gamma\}$ -approximate square.

In the last example we will like to consider  $\delta=1$  and  $\gamma=4$ . This means that we are trying to locate runs of length at least 2 containing only values not greater than  $2\delta=2$  but with  $\gamma\leq 4$ . We obtain:

Position	Square	Roots	$\gamma$
12	(1,1,2,3,1,4)	(2,0,3), (2,1,3)  or  (2,2,3)	4
13	(1,2,3,1,4,5)	(0,3,4), (1,3,4)  or  (2,3,4)	4

#### 5.3 Pseudo-code

Fig. 5 gives the algorithm that solves the  $\delta$ -approximation square problem. Fig. 6 below gives the algorithm that solves the  $\{\delta, \gamma\}$ -approximation square problem.

```
procedure DeltaSquares(t,\,\delta) { n=|t| }
1.
2.
      begin
          for diag \leftarrow 2 to n/2 do
3.
4.
             i \leftarrow 0; dsum \leftarrow 0
              for j \leftarrow diag to n do
5.
6.
                 diff \leftarrow |t[i] - t[j]|
7.
                 if diff \leq \delta then dsum \leftarrow dsum + 1
8.
                  else dsum \leftarrow 0
9.
                 if dsum \ge diag then write j - 2 * diag + 2
10.
                 i \leftarrow i + 1
11.
              od
12.
          od
13. end
```

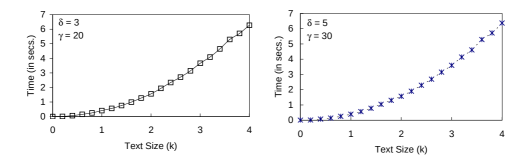
Fig. 5. The DeltaSquares Algorithm.

```
procedure DeltaGammaSquares(t,\,\delta,\,\gamma) { n=|t| }
1.
2.
      begin
3.
          for diag \leftarrow 2 to n/2 do
4.
               \mathbf{i} \leftarrow \mathbf{0}; \; dsum \leftarrow \mathbf{0}; \; gsum \leftarrow \mathbf{0}
               for j \leftarrow diag to n do
5.
                  diff \leftarrow |t[i] - t[j]|
6.
7.
                  if diff \leq \delta then
8.
                  begin
9.
                      dsum \leftarrow dsum + 1
10.
                      gsum \leftarrow gsum + diff
                      if dsum > diag then gsum \leftarrow gsum - |t[i - diag] - t[j - diag]|
11.
12.
                   end
13.
                  else
14.
                  begin
                      dsum \leftarrow 0; gsum \leftarrow 0
15.
16.
17.
                   if dsum \ge diag \text{ AND } gsum \le g \text{ then write } j-2*diag+2
18.
                   i \leftarrow i + 1
19.
               od
20.
           od
21. end
```

Fig. 6. The DeltaGammaSquares Algorithm.

## 5.4 Running time

The complexity of these algorithms is easily seen to be  $O(n^2)$ . Figure 7 shows the timing for different text sizes.



**Fig. 7.** Timing curves for  $\{\delta, \gamma\}$ -approximate squares.

# 6 Conclusion and Open problems

The running time of the computation of  $\delta$ -approximate squares can be reduced to  $O(n \log n)$ ; A theoretical algorithm is presented in [16] that shadows the Main and Lorentz algorithm ([22]).

The following two problems are still open:

**Problem 1.** Given a string  $t = t_1 \dots t_n$  and an integer  $\delta$ , compute all positions j of t, that there exists a string  $\hat{t}$  such that

$$t[j..j+m] \stackrel{\delta}{=} \hat{t}$$
 
$$t[j+m+1..j+2m] \stackrel{\delta}{=} \hat{t}$$
 ... 
$$t[j+\ell m+1..j+(\ell+1)m] \stackrel{\delta}{=} \hat{t}$$

**Problem 2** Given a string  $t = t_1 \dots t_n$  and an integer  $\delta$ , compute all positions j of t, that there exist a string  $\hat{t}$  such that

$$\begin{split} t[j..j+m] &\overset{\delta,\gamma}{=} \hat{t} \\ t[j+m+1..j+2m] &\overset{\delta,\gamma}{=} \hat{t} \\ &\dots \\ t[j+\ell m+1..j+(\ell+1)m] &\overset{\delta,\gamma}{=} \hat{t} \end{split}$$

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Pitch Interval Pattern: {1,-1,3,0}

Pitch Interval String:

 $\{\underline{1,-1,3,0},-5,\underline{2,-2,3,0},-5,0,2,0,2,3,-2,-1,-2,2,\underline{1,-1,3,0},-5,\underline{2,-2,3,0},-5,2,2,1,-1,-2,-2,7,\underline{2,-2,2,0},3,-1,-2,0,-2,0,0,-3,-4,7,-2,-3,5,-3,-4,4,-2,2,\underline{1,-1,3,0},-5,\underline{2,-2,3,0},-5,2,2,1,-1,-2,0,2,0,1,2,2,2,1,-12,4,-2,10,-9\}$ 

#### APPENDIX I

Melody from Mozart's Sonata in A major

This melody may be represented as a string of pitch intervals (in number of semitones). If exact matching is employed, three identical instances of the search pattern are found (patterns a, c and f); the other 4 instances are not matched. If  $\delta$ -approximate matching is employed for  $\delta$ =1, then all seven instances depicted above are found.



Pitch Interval Pattern: {5,-1,1,4,3,5,0}

Pitch Interval String:

 $\{\underline{5,-1,1,4,3,5,0},-1,-2,-2,5,-10,2,1,4,-9,2,2,3,-5,-7,\underline{5,-1,1,4,3,9,0},-2,-2,-1,1,4,-7,3,-1,-1,-1,2,-4,-12,$ 

<u>5,-1,1,4,3,3,0,</u>-1,-2,-2,4,-7,2,1,-1,-2,-5,3,<u>5,-1,1,4,3,5,0,</u>-1,-2,-2,4,-7,2,1,-1,-2,-5,-2,-7,

 $\underline{5,-1,1,4,3,5,0},-1,-2,-2,5,-10,2,1,4,-9,2,2,3,-5,-7,\underline{5,-1,1,4,3,9,0},-2,-2,-3,-2,5,-10,2,1,4,-7,2,1,4,-12,2,1\}$ 

## APPENDIX II

Melody from Schumann's Träumerei

This melody may be represented as a string of pitch intervals (in number of semitones). If exact matching is employed only 3 identical instances of the given pattern are found (patterns a, d and e); the other 3 instances are not matched. If  $\delta$ -approximate matching is employed for  $\delta$ =2, then 4 instances are found (patterns a, c, d and e); for  $\delta$ =4 all 6 instances depicted above are discovered ( $\gamma$ -approximate matching may be additionally applied to restrict  $\delta$ -approximate matching especially for larger  $\delta$  values and for larger melodic corpuses.