Incremental Phylogenetics by Repeated Insertions: An Evolutionary Tree Algorithm

Peter Z. Revesz, Zhiqiang Li

Abstract—We introduce the idea of constructing hypothetical evolutionary trees using an incremental algorithm that inserts species one-by-one into the current evolutionary tree. The method of incremental phylogenetics by repeated insertions lead to an algorithm that can be used on DNA, RNA and amino acid sequences. According to experimental results on both synthetic and biological data, the new algorithm generates more accurate evolutionary trees than the UPGMA and the Neighbor Joining algorithms.

Keywords—Amino acid, DNA, evolution, phylogenetic tree.

I. INTRODUCTION

CURRENT phylogenetic tree construction algorithms[1]-[3], [6], [10], [12] and [14] are not incremental and have to be rerun from the beginning whenever a new species is added to the database. Moreover, a rerun from the beginning is necessary even if the new species is aligned with the already used species. In this paper, we develop an incremental algorithm that inserts new species one-by-one into a growing phylogenetic tree.

Our inspiration for such an incremental phylogenetic algorithm is the way biologists usually classify any newly discovered species. Starting from the root node of the existing classification tree, the newly discovered species is compared with existing species and always an appropriate branch is chosen to go one level down in the classification hierarchy. Eventually we reach one of the existing species, which is the closest relative. It is next to that nearest relative where the new species is normally inserted.

Our aim is to develop a computer algorithm that uses the above paradigm but works with both DNA sequences and proteins. As the genomes of a growing number of species are sequenced and become part of DNA and protein databases [5], [13], molecular biology increasingly augments, although not completely replaces, morphological considerations.

Reliable phylogenetic tree constructions are needed for a diverse set of studies, including theoretical studies on the rate of evolution in various phyla [11] and applied studies aimed at developing medical diagnosis methods [7] and pharmaceutical development. Our algorithm has two main benefits compared to previous algorithms:

1) Faster because it can be used incrementally if the new sequence is aligned with the other sequences.  
2) Generates more accurate phylogenetic trees as indicated by the computer experiments presented in Section 4.

This paper is organized as follows. Section II presents some related work. Section III describes the incremental phylogenetic tree algorithm. Section IV presents some experimental results. Finally Section V gives some conclusions and directions for future work.

II. RELATED WORK

The UPGMA [11] and the Neighbor Joining (NJ) [10] algorithms are commonly used and familiar to most users. The maximum likelihood method is also well known, although it seems less frequently used that UPGMA and Neighbor Joining in practice because it requires more computational time. All of these algorithms are reviewed in textbooks, such as [1]-[3].

Revesz [6] introduced the Common Mutations Similarity Matrix algorithm, which has O(n³) time complexity, where n is the number of sequences. We briefly review this algorithm as a related work, which will also be used in the experimental results section of this paper. Table I below shows seven DNA sequences, S₁…S₇, each with a length fifteen nucleotides displayed by groups of five nucleotides per column.

| S1  | AGCTA | CTAGT | AATCA |
| S2  | AGCTA | CGAGT | AATCA |
| S3  | ATCCA | CTAGT | ACACT |
| S4  | ATCCA | CTAGT | ATACT |
| S5  | CGTCA | TTGTA | AAGCT |
| S6  | CGTCA | CATCA | AATGC |
| S7  | AGGTA | CTGGA | AATCC |
| μ   | AGCTA | CTAGT | AATCT |

Let S_i[k] denote the kth nucleotide of Si. The Hamming distance between two DNA sequences S_i and S_j each with length n, denoted δ (S_i, S_j), is defined as the number of corresponding nucleotide pairs that are different, that is, \( \Sigma l \leq k \leq n S_i[k] \neq S_j[k] \). μ is the common ancestor of seven sequences.

Evolutionary tree construction algorithms generally start

Peter Z. Revesz is with the Department of Computer Science and Engineering, University of Nebraska-Lincoln, Lincoln, NE 68588, USA (revesz@cse.unl.edu).
Zhiqiang Li is with the Department of Computer Science and Engineering, University of Nebraska-Lincoln, Lincoln, NE 68588, USA (zli@cse.unl.edu).
from a Hamming distance matrix to recursively combine pairs of sequences (rows and columns) until only a single combined sequence remains. For example, the UPGMA (unweighted pair group method with arithmetic mean) [12] method would always search for the closest pairs to combine. When several pairs are equally distant, then an arbitrary choice is made. In this case, the closest pairs are S1 and S2 and S3 and S4 because \( \delta (S1, S2) = 1 \) and \( \delta (S3, S4) = 1 \). The Neighbor Joining [10] method is a more sophisticated and commonly used method that is also based on distance matrices.

Instead of distance matrices, Revesz [6] introduced a common mutations similarity matrix (CMSM). The motivation behind looking for common mutations is that in practice rare but shared features, such as rare mutations, often provide useful markers of similarity among a set of closely related items. Moreover, if mutations are rare, then it may be more efficient to count their occurrences than finding the Hamming distances for long sequences. Assuming that the seven DNA sequences in Table 1 are related, we can find the most likely common ancestor sequence, denoted \( \mu \), as the mode of each column. If there is no most frequent nucleotide in a column, then we arbitrarily chose one of the most frequent nucleotides in it.

The Common Mutations Similarity Matrix (CMSM) algorithm records for each pair of sequences the mutations that they share in common with respect to a global average \( \mu \), which is taken as the most likely common ancestor sequence.

**Example 1.** Given seven nucleotide sequences in Table 1 below (rows S1 to S7 where the sequences are displayed in groups of five), the common ancestor sequence \( \mu \) is calculated in [6] as the most frequent in each column.

Alternatively, if S1 to S7 are considered amino acid sequences where A, C, G and T now stand for the amino acids Alanine, Cysteine, Glycine and Threonine, respectively, then the common ancestor sequence \( \mu \) can be defined as in each column as the amino acid \( x \) out of the set \( S \) of twenty amino acids used in most proteins such that \( x \) is overall closest to the set of amino acids in that column. We make this statement more precise below using as an example the PAM250 amino acid similarity matrix. Let

\[
\text{PAM250}(\text{AminoAcid}1, \text{AminoAcid}2) = a
\]

denotes that AminoAcid1 and AminoAcid2 have a similarity score of \( a \). For example, PAM250 \([A, G] = 1\) means that Alanine and Glycine are slightly similar to each other. Then for the \( i \)th column,

\[
\mu[i] = x \in S
\]

such that

\[
\sum_{j=1}^{n} \text{PAM250}[S_i[j], x]
\]

is maximum.

For example, we can see that the value of \( \mu[1] \) changed from A to C because C is the amino acid that is overall closest to the each of the amino acids in the first column.

<table>
<thead>
<tr>
<th>S1</th>
<th>S2</th>
<th>S3</th>
<th>S4</th>
<th>S5</th>
<th>S6</th>
<th>S7</th>
</tr>
</thead>
<tbody>
<tr>
<td>AGCTA</td>
<td>CTAGT</td>
<td>AATCA</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AGCTA</td>
<td>CGAGT</td>
<td>AATCA</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ATCCG</td>
<td>CTAGT</td>
<td>ACACT</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ATCCG</td>
<td>CTAGT</td>
<td>ATACT</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CGGTA</td>
<td>TTGTG</td>
<td>AAGCT</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CGGTT</td>
<td>CATCA</td>
<td>AATGC</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AGGTA</td>
<td>CTGTA</td>
<td>AATCC</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CGCCA</td>
<td>TTTGT</td>
<td>AATCC</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

It can be assumed that in each sequence Si those amino acids (or nucleotides) that do not match the corresponding amino acid (or nucleotide) in \( \mu \) were mutated at some point during evolution. Intuitively, the more common mutations two sequences \( S_i \) and \( S_j \) share, the closer they are likely to be in an evolutionary tree. For the above set of sequences, the common mutations similarity matrix is shown in Table 3:

<table>
<thead>
<tr>
<th>S1</th>
<th>S2</th>
<th>S3</th>
<th>S4</th>
<th>S5</th>
<th>S6</th>
<th>S7</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>4</td>
<td>2</td>
<td>2</td>
<td>1</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>4</td>
<td>0</td>
<td>2</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>0</td>
<td>5</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>5</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>1</td>
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<tr>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>2</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>0</td>
</tr>
</tbody>
</table>

According to the common mutations similarity matrix, the closest pair of sequences is S3 and S4. Hence these will be merged. When we merge two sequences \( S_i \) and \( S_j \), in the merged sequence the \( k \)th element will be equal to the amino acid (or nucleotide) in the two sequences if \( S_i[k] = S_j[k] \) and will be equal to \( \mu[k] \) otherwise. Hence the matrix of sequences will be updated as Table 4:

<table>
<thead>
<tr>
<th>S1</th>
<th>S2</th>
<th>S3</th>
<th>S4</th>
<th>S5</th>
<th>S6</th>
<th>S7</th>
</tr>
</thead>
<tbody>
<tr>
<td>AGCTA</td>
<td>CTAGT</td>
<td>AATCA</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AGCTA</td>
<td>CGAGT</td>
<td>AATCA</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ATCCG</td>
<td>CTAGT</td>
<td>AAACCT</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CGGTA</td>
<td>TTGTG</td>
<td>AAGCT</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CGGTT</td>
<td>CATCA</td>
<td>AATGC</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AGGTA</td>
<td>CTGTA</td>
<td>AATCC</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CGCCA</td>
<td>TTTGT</td>
<td>AATCC</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

For example, since \( S3[12] = C \neq T = S4[12] \), by the above merging rule \( S34[12] = \mu[12] = A \).

After the merge, the common mutations matrix needs to be recalculated. The merge does not change \( \mu \), but the entries in the common mutations similarity matrix that are related to the newly merged sequence S34 need to be calculated. The values for S3 and S4 should be deleted. In this case, Table 5 shows the updated common mutation matrix.
Table 5 The updated CMSM matrix

<table>
<thead>
<tr>
<th></th>
<th>S1</th>
<th>S2</th>
<th>S3</th>
<th>S4</th>
<th>S5</th>
<th>S6</th>
<th>S7</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1</td>
<td>0</td>
<td>4</td>
<td>2</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>S2</td>
<td>4</td>
<td>0</td>
<td>2</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>S34</td>
<td>2</td>
<td>2</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>S5</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>2</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>S6</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>2</td>
<td>0</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>S7</td>
<td>2</td>
<td>2</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>0</td>
<td></td>
</tr>
</tbody>
</table>

Now the closest pair is S1 and S2 with a value of 4 common mutations. Hence those two will be merged next. The merging will continue until there is only one sequence left. The CMSM evolutionary tree algorithm can be summarized as shown in Fig. 1.

Algorithm CMSM (S1...Sn, n)
1. Form n clusters of sequences, each with a single sequence.
2. Find the putative common ancestor µ of the sequences.
3. Construct a graph T with a node for each n cluster and for µ.
4. While (there is more than one cluster)
   5. Find the common mutations similarity matrix.
   6. If (exist distinct Si and Sj with some common mutations)
      7. Merge a closest distinct Si and Sj pair into a new cluster Sij and create a node for Sij.
      8. Connect the nodes for Si and Sj with parent node Sij.
   9. Else
      10. Connect the remaining clusters' nodes to parent µ.
      11. Return T.
12. Return T.

Fig. 1 The CMSM algorithm

Note: Alternatively, instead of only recording the values, the actual set of common mutations can be put into each entry of the common mutations similarity matrix. Clearly, the cardinality of the sets in the second representation determines the numerical values in the first representation.

III. INCREMENTAL PHYLGENETICS BY REPEATED INSERTIONS

A. A New Phylogenetic Tree Algorithm

Suppose that we have n number of amino acid sequences S1, . . ., Sn. The sequences and the number n are the inputs to the following algorithm that constructs an evolutionary tree by repeated addition of new species that are represented by the amino acid sequences. We call the new algorithm IPRI (incremental phylogenetic by repeated insertions). A pseudo-code of the IPRI algorithm is shown in Fig. 2.

In the algorithm, the closest pair can be found by minimum Hamming distance if the sequences are DNA or RNA strings. If the sequences are proteins, then the closest pair can be found by using a PAM or a BLOSUM substitution matrix. The running time is O(n^2m) where m is the length of the sequences because there are n insertions, and each insertion requires n comparisons between two strings of length m.

Algorithm IPRI(S1...Sn, n)
1. Create an independent node Nk for each sequence Sk.
2. Let N = \{ Nk : 1 \leq k \leq n \}
3. Find the closest pair of nodes Nj and Nj.
4. Create a tree T with root R, left child Ni and right child Nj.
5. N = N \setminus \{ Ni, Nj \}
6. While (N is not empty)
   7. Find the closest pair of nodes Ni \in N and Mj \in T.
   8. If (Mj is not the root of T)
     10. Delete P as a parent of Mj.
     11. Create a node R.
     12. Make P the parent of R.
     13. Make R the parent of Ni and Mj.
     14. Else
        15. Create a node R.
        16. Make R the parent of Ni and Mj.
     17. N = N \setminus \{ Ni \}.
18. Return T.

Fig. 2 The IPRI algorithm

The two major cases of insertion in the IPRI algorithm are illustrated in Fig. 3 and Fig. 4, respectively.

Fig. 3 shows the case when the new node to be inserted, node 3, is closest to the root node R. In this case, the IPRI algorithm creates a new root called P and makes both the old root R and the newly inserted node 3 the children of P.

Fig. 4 shows the case when the new node, again numbered node 3, is closest to node 2. In this case, the IPRI algorithm creates a new node P, which becomes a child of R, while both nodes 2 and 3 become children of P. The case when node 3 is closest to node 1 is a symmetric case, which is not illustrated separately.
IV. EXPERIMENTAL RESULTS

A. Experiments with Simulated Data

We compared the algorithms on simulated evolutionary data as follows. We assumed that the original protein consists of a chain of one thousand Alanine amino acids. We mutated this original string two ways to generate to children. Both children were generated by first randomly selecting one percent of the amino acids. Then we changed the selected amino acids to one of the twenty amino acids. That is, each of the selected amino acids had a five percent chance of remaining A and ninety five percent chance of changing into another amino acid, with five percent chance of changing into C, five percent chance of changing into D and so on.

Next both of the children were further mutated to generate four grandchildren of the original protein. Then we generated additional levels of the tree so that after N levels we had \(2^N\) leaves.

With the above process of evolutionary tree generation, two siblings can be expected to differ from each other on twenty amino acids. Two first cousins can be expected to differ from each other on forty amino acids. Two seconds can be expected to differ from each other on sixty amino acids, and so on.

We ran ten tests on evolutionary trees with height four (and sixteen leaves). We implemented the CMSM and the IPRA algorithms in MATLAB. We used ClustalW2's implementation of the UPGMA and NJ algorithms. We chose on the ClustalW2 website the default parameters, that is, a gap open penalty of 10, a gap extension penalty of 0.2, and a maximum gap distance of 5. The results can be summarized in the Table 8, where ``Perfect'' means that the reconstructed tree is the same as the original evolutionary tree. When a reconstructed tree had errors, we checked only how many of the sibling pairs (SPs) were correctly handled.

### Table 8 Experimental comparisons of the algorithms

<table>
<thead>
<tr>
<th>Test</th>
<th>CMSM</th>
<th>IPRA</th>
<th>UPGMA</th>
<th>NJ</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Perfect</td>
<td>Perfect</td>
<td>8 SPs</td>
<td>8 SPs</td>
</tr>
<tr>
<td>2</td>
<td>Perfect</td>
<td>Perfect</td>
<td>7 SPs</td>
<td>7 SPs</td>
</tr>
<tr>
<td>3</td>
<td>Perfect</td>
<td>Perfect</td>
<td>7 SPs</td>
<td>7 SPs</td>
</tr>
<tr>
<td>4</td>
<td>Perfect</td>
<td>Perfect</td>
<td>6 SPs</td>
<td>7 SPs</td>
</tr>
<tr>
<td>5</td>
<td>Perfect</td>
<td>Perfect</td>
<td>7 SPs</td>
<td>7 SPs</td>
</tr>
<tr>
<td>6</td>
<td>Perfect</td>
<td>Perfect</td>
<td>7 SPs</td>
<td>7 SPs</td>
</tr>
<tr>
<td>7</td>
<td>Perfect</td>
<td>Perfect</td>
<td>8 SPs</td>
<td>8 SPs</td>
</tr>
<tr>
<td>8</td>
<td>Perfect</td>
<td>Perfect</td>
<td>8 SPs</td>
<td>8 SPs</td>
</tr>
<tr>
<td>9</td>
<td>Perfect</td>
<td>Perfect</td>
<td>6 SPs</td>
<td>6 SPs</td>
</tr>
<tr>
<td>10</td>
<td>Perfect</td>
<td>Perfect</td>
<td>7 SPs</td>
<td>7 SPs</td>
</tr>
</tbody>
</table>

As an example, Fig. 5 shows the output of the IPRA algorithm in case 4. As a comparison, Fig. 6 shows the output of the UPGMA algorithm in the same case.

B. Experiments with Biological Data

In this section, we describe experiments with both telomerase protein and telomerase RNA data.

### Telomerase Protein Experiments:

We investigated the telomerase (TERT) protein family. Telomerase help protect eukaryote chromosomes during duplication. From the website http://telomerase.asu.edu we obtained 14 vertebrate telomerase proteins as our input data. After alignment, the length of each amino acid sequence was 1353.

![Fig. 5 Sample evolutionary tree reconstructed by the IPRA algorithm](image)

The IPRI algorithm, which we implemented in MATLAB, with the gap penalty value -1, gave the phylogenetic tree shown in Fig. 7. Using ClustalW2 with gap penalty 10 and gap extension 0.1 and the same telomerase RNA input data we also generated the UPGMA and the Neighbor Joining phylogenetic trees, which are shown in Fig. 8 and Fig. 9, respectively. We also implemented the CMSM algorithm in MATLAB. The CMSM phylogenetic tree is shown in Fig. 10.

### Telomerase RNA Experiments:

We also investigated the telomerase RNA (TR) family. From the website http://telomerase.asu.edu we obtained 42 vertebrate telomerase RNA as our input data. After alignment, the length of each RNA sequence was 741.

The IPRI algorithm gave the phylogenetic tree shown in Fig. 11. Using ClustalW2 and the same telomerase input data we obtained the UPGMA and the Neighbor Joining phylogenetic trees shown in Fig. 12 and Fig. 13, respectively. Finally, Fig. 14 shows the CMSM phylogenetic tree.
Fig. 7 The IPRI phylogenetic tree based on vertebrate telomerase protein data
Fig. 8 The UPGMA phylogenetic tree based on vertebrate telomerase protein data

Fig. 9 The Neighbor Joining phylogenetic tree based on vertebrate telomerase protein data

Fig. 10 The CMSM phylogenetic tree based on vertebrate telomerase protein data
Fig. 11 The IPRI phylogenetic tree based on vertebrate telomerase RNA data
Fig. 12 The UPGMA phylogenetic tree based on vertebrate telomerase RNA data
Fig. 13 The NJ phylogenetic tree based on vertebrate telomerase RNA data
Fig. 14 The CMSM phylogenetic tree based on vertebrate telomerase RNA data
C. Discussion of the Experimental Results

We divide the discussion into three parts. In the first part we discuss the simulated data results, in the second part the protein results, and in the third part the RNA results.

1. Simulated Data Results: The simulated data results suggested that the new IPRA algorithm is an improvement over the older UPGMA algorithm. For example, as can be seen from Fig. 5 and Fig. 6, the IPRA algorithm has given back the original evolutionary tree in both cases. On the other hand, the UPGMA algorithm made a mistake in some of the sibling pairs. In particular, the leaves 26 and 27 and the leaves 18 and 19 are not paired correctly. In addition, there are more mistakes in grouping together cousin leaves. For example, the sibling leaves 16 and 17 are paired correctly, but they are not grouped correctly with their cousin leaves 18 and 19.

2. Telomerase Protein (TERT) Results: The UPGMA and the Neighbor Joining phylogenetic trees suppose that vertebrate evolution started with the mammals. According to Fig. 8 and Fig. 9, the mammals started to diverge early on and all the other non-mammal vertebrates are like one small branch of the big mammalian evolutionary tree. In contrast, the IPRI phylogenetic tree in Fig. 7 separates the mammals and the non-mammals into two parallel branches. The CMSM tree in Fig. 10 has the fish branch out first, then the reptiles and the birds and finally the mammals. Hence the IPRI and the CMSM phylogenetic trees are more realistic. However, it is possible that the UPGMA and the Neighbor Joining results would improve if we considered a larger set of proteins from the same protein family.

3. Telomerase RNA (TR) Results: In this case, as shown in Fig. 11, the IPRI algorithm followed well the accepted evolutionary theory. In the IPRI algorithm, the fish is the earliest vertebrate group that separates from the other vertebrates, followed by the amphibians. The subtree with root 68 consists of all the mammals. In contrast, the UPGMA and the Neighbor Joining phylogenetic trees still make the mistake of assuming that mammals were the earliest vertebrate group. Therefore the UPGMA and the Neighbor Joining phylogenetic trees run completely counter to the accepted order of vertebrate evolutionary history. Finally, the CMSM result was also unrealistic because, for example, it put together in the subtree rooted at node 75 some fish and various mice.

V. CONCLUSIONS AND FUTURE WORK

The new incremental phylogenetic tree algorithm has a potential to improve the general phylogenetic trees and our understanding of evolutionary history, as can be inferred based on molecular biology. Generally, all phylogenetic tree algorithms improve with greater data size both with the number of species and in the length of the sequences. In the future, we plan to study additional protein families and their DNA and amino acid sequences. Finally, it would be interesting to look at the evolution of biological vision in order to learn from it ideas that may improve digital cameras [4].

REFERENCES


Peter Z. Revesz holds a Ph.D. degree in Computer Science from Brown University. He was a postdoctoral fellow at the University of Toronto before joining the University of Nebraska-Lincoln, where he is a professor in the Department of Computer Science and Engineering. Dr. Revesz is an expert in bioinformatics, databases, data mining, and big data analytics. He is the author of Introduction to Databases: From Biological to Spatio-Temporal (Springer, 2010). Dr. Revesz held visiting appointments at the IBM T. J. Watson Research Center, INRIA, the Max Planck Institute for Computer Science, the University of Athens, the University of Hasselt, the U.S. Air Force Office of Scientific Research and the U.S. Department of State. He is a recipient of an AAAS Science & Technology Policy Fellowship, a J. William Fulbright Scholarship, an Alexander von Humboldt Research Fellowship, a Jefferson Science Fellowship, and a National Science Foundation CAREER award.

Zhiqiang Li got his B.S. and M.S. degrees from Xidian University, China, in 2009 and 2012, respectively. He is currently a Ph.D. student in the University of Nebraska-Lincoln. His research interests include bioinformatics, computer security, data mining and image processing.