



IN APPLIED SCIENCE & ENGINEERING TECHNOLOGY

Volume: 8 Issue: VI Month of publication: June 2020

DOI: http://doi.org/10.22214/ijraset.2020.6112

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Application of Visual Basic in Bioinformatics for the Structural Prediction of Protein

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Abstract: Protein secondary structure prediction is a problem of central importance in predictive structural biology. It has been studied intensively for more than 30 years. However, the best models currently available to perform this task suffer from a fundamental drawback their prediction is only based on the content of a small segment of the sequence surrounding the residue the conformational state of which is to be determined when compared with the database searches it is good to use programming language (Visual Basic) The choufasman algorithm developed using visual basic for the secondary structure of protein plays a significant role. The protein structure prediction using choufasman in visual basic showed prediction accuracy of 56 to 60%. In these studies the protein catalase was selected and using the visual basic programming language the amino acid sequence for protein catalase it's secondary structure and choufasman algorithm with score was also analyzed.

Keywords: Catalase, secondary structure prediction, visual basic, chou Fasman algorithm, SOPMA..

I.

A. What is Bioinformatics?

Bioinformatics is a field that develops database, methods and software tools for studying biological data, when the data sets are large and complex Bioinformatics involves the integration of computers, software tools, and databases. Bioinformatics approaches are often used for major initiatives that generate large data sets. The genomics and proteomics are the two important collection, of large-scale activities that use bioinformatics. Classification, storage, and analysis of biochemical and biological information using computers especially as applied to molecular genetics and genomics. Bioinformatics has become an important part of many areas of biology. It plays an important role in structural prediction of protein, genome analysis, DNA and RNA binding region, Homology modeling, and identification of drug compounds. It plays a role in the analysis of gene and protein expression and regulation.

INTRODUCTION

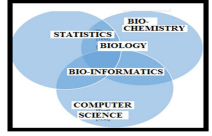


Figure1 Bio-informatics overview

B. Visual Basic

Visual Basic (VB) is a third generation event-driven programming language first released by Microsoft in 1991.Microsoft that provides a graphical user interface (GUI) which allows programmers to modify code by simply dragging and dropping objects and defining their behavior and appearance. Visual Basic can create executable (EXE files), ActiveX controls, or DLL files, but is primarily used to develop Windows applications and to interface database systems. Dialog boxes with less functionality can be used to provide pop-up capabilities. NET (VB.NET) is a general-purpose programming language that works in the NET environment.



Figure 2 Visual Basic



ISSN: 2321-9653; IC Value: 45.98; SJ Impact Factor: 7.429 Volume 8 Issue VI June 2020- Available at www.ijraset.com

C. Structural Prediction Of Protein

Proteins are polymers of amino acids covalently linked through peptide bonds into a chain. They constitute a group of highly complex organic compounds found in all living cells and comprising the most abundant class of all biological molecules. Protein comprises approximately 50% of cellular dry weight. Hundreds of protein molecules have been isolated in pure, homogeneous form; many have been crystallized.

All contain carbon, hydrogen, and Oxygen, and nearly all contain sulfur as well. Some proteins also incorporate phosphorous, ion, zinc, and copper.

Proteins are large molecules with high molecular weights from about 10,000 for small ones of 50-100 amino acids to more than 1,000,000 for certain forms.

They are composed of varying amounts of the same 20 acids, which in the intact protein are united through covalent chemical linkages called peptide bonds.

Within and outside of cells, proteins serve a myriad of functions, including structural roles (cytoskeleton), as catalysts (enzymes transporter to ferry ions and molecules across membranes, and hormones.

A protein molecule that consists of but a single polypeptide chain is said to be monomeric; proteins made up of more than one polypeptide chain, as many of the large ones are called oligomer.

Based upon chemical composition, proteins are divided into two major classes: simple proteins, which are composed of only amino acids, and conjugated proteins which are composed of amino acids and additional organic and inorganic groupings, certain of which are called prosthetic groups. Conjugated proteins include glycoproteins, which contain carbohydrates; lipoproteins, which contain lipids; and nucleoproteins, which contain nucleic acid

Classified by biological function, proteins include the enzyme, which are responsible for catalyzing the thousands of chemical reactions of the living cell; keratin, elastin, and collagen, which are important types of structural, or support, proteins; hemoglobin and other gas transport proteins; ovalbumin, casein, and other nutrient molecules; antibodies, which are molecules of the immune system; protein hormones, which regulate metabolism : and proteins that perform mechanical work, such as actin and myosin, the contractile muscle proteins

D. Classes Of Protein Structure

- 1) Class Alpha- this comprises a bundle of alpha helices connected by loops on the surface of the proteins.
- 2) Class Beta- this comprises antiparallel beta sheets, usually two sheets in close contact forming a sandwich. Alternatively, a sheet can twist in to a barrel with the first and last strands touching. Examples are enzymes, transport proteins, antibodies and virus coat proteins such as neuraminidase.
- 3) Class Alpha/Beta- this comprises mainly parallel beta sheets with intervening alpha helices, but may also have mixed beta sheets. In addition to forming a sheet in some proteins in this class, in others parallel beta strands may form in to a barrel structure that is surrounded by alpha helices. This class of proteins includes many metabolic enzymes.
- 4) Class Alpha+Beta- this comprises mainly segregated alpha helices and antiparallel beta sheets.
- 5) Multidomain (Alpha and Beta)- Proteins comprise domains representing more than one of the above four classes.
- 6) Membrane and cell surface- Proteins and peptides excluding proteins of the immune system comprise this class.

E. Levels Of Protein Structure

Structural features of proteins are usually described at four levels of complexity

- 1) *Primary Structure:* The linear arrangement of amino acids in a protein and the location of covalent linkages such as disulfide bonds between amino acids.
- 2) Secondary Structure: The areas of folding or coiling within a protein; examples include alpha helices and pleated sheets, which are stabilized by hydrogen bonding.
- 3) *Tertiary structure:* The final three-dimensional structure of a protein, which results from a large number of non-covalent interaction between amino acids.
- 4) *Quaternary Structure:* Non-covalent interactions that bind multiple polypeptides into a single, larger protein. Hemoglobin has quaternary structure due to association of two alpha globin and two beta globin polyproteins.



International Journal for Research in Applied Science & Engineering Technology (IJRASET)

ISSN: 2321-9653; IC Value: 45.98; SJ Impact Factor: 7.429 Volume 8 Issue VI June 2020- Available at www.ijraset.com

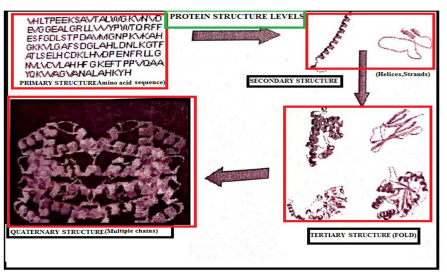


Figure3 protein structure types

II. REVIEW OF PRIMARY STRUCTURE OF PROTEIN AND TERMINOLOGY

Proteins are polymers of amino acids joined together by peptide bonds. The Primary structure of a protein can readily be deduced from the nucleotide sequence of the corresponding messenger RNA. Based on primary structure, many features of secondary structure can be predicted with the aid of computer programs. However, predicting protein tertiary structure remains a very tough problem, although some progress has been made

A. Assembly of Primary Structure

Synthesizing primary structure of a protein is encoded in genetic material of the chromosomes, which is double stranded DNA in most organisms though some viruses use single stranded DNA or RNA. In all cases the information is coded as sequences of 4 kinds of nucleotides on one strand of DNA - A, C, G, and T or of RNA which has the same the nucleotides except U which replaces T. The sequence of other strand of DNA is chemically complimentary to first i.e. A with T of other strand and G with C.

Specific region of DNA or RNA codes for primary structure of every protein synthesized by organism and the DNA, RNA segments on both sides of such regions are involved in regulation and expression of that genetic information, these regions of DNA or RNA are called Genes. Parts of a gene that code for primary structures of protein are well defined but regulatory parts are not. The information in nucleic acid sequence of gene is invariably read in one direction from 5' end to 3 end. So it is possible to describe gene structure in terms of upstream and downstream in coding sequence.

Some genes code only for stable RNA molecules such as ribosomal and transfer RNA. Initial steps in expression of genes are same as in transcription into RNA of genes that code for proteins but resulting RNA is not translated into protein.

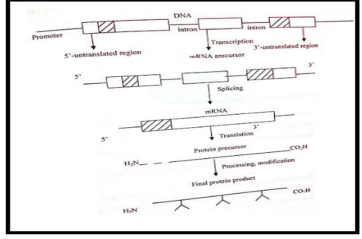
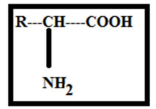


Figure 4 Assembly of protein structure



B. Amino Acids

Amino acids appear in all proteins in all forms of life (and viruses). Amino acids "condense" in the ribosome to form proteins Amino acids are the building blocks of protein. The general structure of an amino acid is shown below.



Each amino acid is a nitrogenous compound consisting of an acidic carboxyl, COOH) group and a basic amino, (- NH2) group. To the alpha carbon, a side chain (often denoted as - R) is attached. R can be simple as a hydrogen atom (H) or a methyl group CH2) or a more complex structure. The side chains vary with each amino acid, and these various side chains confer unique stereo chemical properties on each amino acid.

C. Secondary Structure

Protein structure plays an important role in its function. The protein structure may not be functional if it loses its structure. Primary structure is the amino acid sequence. Secondary structure is local interactions between stretches of a polypeptide chain and includes α -helix and β -pleated sheet structures. Stretches or strands of proteins or peptides have distinct, characteristic local structural conformations, or secondary structure, dependent on hydrogen bonding. The two main types of secondary structure are the α -helix and the β -sheet. The α -helix is a right-handed coiled strand. In common there are three types of secondary structures in proteins, namely alpha helices, beta sheets, and turns. That which cannot be classified as one of the standard three classes is usually grouped into a category called "other" or "random coil".

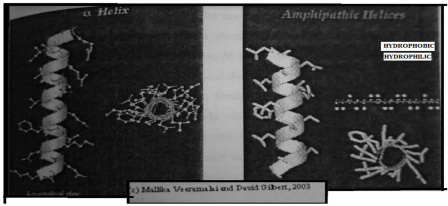


Figure5 Secondary Structure

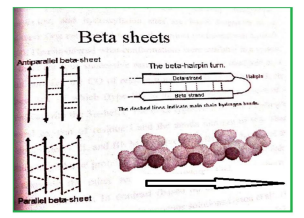


Figure 6 Beta-sheets



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D. Tertiary Structure

Protein tertiary structure is the three dimensional shape of a protein. The tertiary structure will have a single polypeptide chain "backbone" with one or more protein secondary structures, the protein domains. Amino acid side chains may interact and bond in a number of ways (Wikipedia)

Tertiary structure is the next level of complexity in protein folding the tertiary structure of proteins deals with how the regional structures are put together in space (www.sciencedirect.com).

E. Quaternary Structure

The quaternary structure of a protein was usually determined by x-ray crystallography .It is the association of several protein chains and subunits into a closely packed arrangement. The protein subunits has its own primary, secondary, and tertiary structure. Protein subunits are held together by hydrogen bonds and van der Waals forces between nonpolar side chains.

III. USING VISUAL BASIC FOR PROTEIN STRUCTURE PREDICTION

A. Program 1(Amino acid Sequence, mRNA sequence prediction) Private Sub MDIForm_Load () End Sub Private Sub mnuAmino ClickO IFlag "MAIN" FrmAmino.Show End Sub Private Sub mnuBack_ Click () IFlag= "MAIN" FrmBackProcess.Show End Sub Private Sub mnuExit Click () End End Sub Private Sub mnuMRna Click () IFlag= "MAIN" FrmMrna.Show End Sub Private Sub mnuSec Click () FrmSecondary.Show End Sub _____ B. Program 2 (Secondary Structure Prediction) Dim i as Integer Dim mRNA as String _____ Private Sub Command1 Click () Unload Me End Sub _____ Private Sub Form Load () Me. Width 3735



Me.Height 4455 Me.Top=0 Me.Left= 0 I=0 MRna = "UCAG" Randomize 3 End Sub Private Sub Form Resize () On Error Resume Next txtMrna.Height = Me.ScaleHeight - 500 TxtMrna. Width = Me.Scale Width End Sub

Private Sub mnuCopy_Click () Clipboard.SetText (txtMrna.SelText) End Sub Private Sub mnuCut Click () Clipboard.SetText (txtMrna.SelText) txtMrna.SelText = "" End Sub

Private Sub mnuExit Click () Unload Me

End Sub Private Sub MnuFindAmino_Click () Dim I as Integer |= Len (txtMrna.Text) IFlag = "MRNA" If1< 100 Then MsgBox "MRNA less than 100" Else FrmAmino.Show End If End Sub

Private Sub mnuNew _Click () 0=! TxtMrna. Text = w End Sub

Private Sub mnuOpen_Click () On Error Resume Next cd1.ShowOpen txtMrna.LoadFile (cd1.FileName) MnuFindAmino.Enabled True End Sub

Private Sub mnuPaste_Click () txtMrna.SelStart = Clipboard.GetData () End Sub



Private Sub mnuRandom Click () Dim I, j, limit As Integer Dim s as String TxtMrna. Text = Limit = Input Box ("Input the limit") For i = 1 to limit |= Rnd * 3 I + 1 = 1s = Mid (mRNA, 1, 1)txtMrna.Text = txtMrna.Text & s Next End Sub Private Sub mnuSave Click () On Error Resume Next cd1.ShowSave txtMrna.SaveFile (cd1.FileName), rtfCFText End Sub Private Sub txtMrna Key Down (Key Code as Integer, Shift As If Key Code 65 and Key Code >97 and Key Code 67 and Key code<>99 and

Key Code 05 and Key Code 257 and Key Code 07 and Key Code 259 and Key Code 85 and Key Code 117 and Key code<>71 And Key Code 103 and Key Code<> 8 Then Key Code= 0 End If End Sub

Private sub txtMrna_KeyPress (KeyAscii As integer)

End sub

Private Sub txtMrna_LostFocus ()

End sub

C. Program 3(Back Translation Process) Dim rs As Recordset Dim frame (5) As String Dim strAmino As String Public get Amino as String Dim LOSOURCESTR as String Dim i as Integer

Private Function FindStartCodon (mRNA as String) As String Dim j, I as Integer Dim STR, StrStartCodon As String For j 1 to Len (mRNA) STR = Mid (mRNA, j, 3) If str "AUG" Then



StrStartCodon = str = True Exit For End If Next j 1 = Len (mRNA) -j-2 If1>= 0 Then StrStartCodon = StrStartCodon & Mid (mRNA, j+3, 1) StrStartCodon = StrStartCodon & Mid (mRNA, 1,j-1) FindStartCodon= StrStartCodon Else MsgBox "No start Codon found" Unload Me End If

End Function

Private Sub SubFindFilling (By Val SOURCESTR as String, By Val j as Integer, By Val No as Integer For i=j to Len (SOURCESTR) Step 3 s= Mid (SOURCESTR, i, 3) If Len(s) 3 Then Set rs = db.OpenRecordset ("Select Amino from tblmRNA where Codon=" &s If rs.RecordCount > 0 Then Select Case rs. Fields (0) Case "M" Frame (no) = frame (no) & rs. Fields (0) StrAmino = strAmino & "M" Text show. Text = txtshow.Text & "M" Case "*", "\$", Frame (no) = frame (no) & rs. Fields (0) StrAmino = strAmino & rs. Fields (0)Case Else Frame (no) = frame (no) & rs. Fields (0) StrAmino = strAmino & rs. Fields (0)Text show. Text text show. Text & rs. Fields (0) End Select End If End If PB.Value = PB.Value + 1Next i Text show.Text text show.Text & Chr (13) & Chr (10) StrAmino = strAmino & ";" End Sub

Private Sub Filling (SOURCESTR as String) Dim revstr, desstr As String Dim i, j, k as Integer



Dim s as String For j =0 To 5 Frame (j) ='"" Next strAmino=" "

PB.Value 0 txtshow.Text = ""

For j =1 To 3 Call SubFindFilling (SOURCESTR, j, j - 1) Next

For j Len (SOURCESTR) To 1 Step -1 Revstr = revstr & Mid (SOURCESTR, j, 1) Next j

For j 1 To 3 Call SubFindFilling (revstr, j, j+ 2)

Next j

PB.Visible= False End Sub

Private Sub mnuSave_Click () cd1.ShowSave Open cdl.FileName For Output As #2 Print #2, strAmino Close #2 End Sub

Private Sub mnuSec_Click () Get Amino = strAmino FrmSecondary.Show End Sub %3D

Private Sub txtshow_Change () End Sub

Dim frame (5) As String Dim Amino Acid as String Dim iApa parti1 As Long A Count Partl. A Fcount Partil, A BCount Partil As Long Dim iFApa_ part2, iBApa part2 As Long Dim Alpha Range (100, 1) As Long Dim iBpa_ part! As Long International Journal for Research in Applied Science & Engineering Technology (IJRASET) ISSN: 2321-9653; IC Value: 45.98; SJ Impact Factor: 7.429

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Dim B _Count Parti, B _ FCount Partll, B _ BCount Partll as Long Dim Beta Range (100, 1) As Long Dim Alpha Helices, Beta Sheets as Long Dim Alpha Helices no, Beta Sheets no As Integer Dim Beta Turns As Double Dim Beta Stop Codon as Integer Dim From Where As Long Dim Where Up To As Long Dim ARange_ Count As Long Dim BRange_ Count As Long

Private Sub FindAlphaHelicesPartl () Dim IPA value (5), c, i, j, k, pos As Long Dim countless 100 As Long Dim rs As Recordset Dim flag As Boolean Dim strAnAmino As String From Where = 1While from Where < Len (Amino Acid) StrAnAmino- Mid (Amino Acid, From Where, 1) Flag True Set rs = db.OpenRecordset ("Select pa from Amino Acid where Amino acid=" & strAnAmino & ") If rs.RecordCount >0 Then I + sod = sod $ipa_value(j) = rs.$ Fields (0) Count less100 =0 For k 0 To 5 If ipa value (k) ≤ 100 Or From Where ≤ 5 Then count_less100 = count l-If count less 100 > 2 Then Exit For End If Next k If flag true then Call ForwardFindAlphalHelicesPartIII (Alpha_part1, i) From Where i Exit For Else If> 3 Then For k 0 To 2 $ipa_value(k) = ipa_value(k + 1)$ Next k i=2End If End if J=j+1End if Next i



```
J=0
For i=0 To 3
ipa_value (i) = 0
Next i
For i = (Alpha_partl - 6) To WhereUpTo Step -1
StrAnAmino = Mid (Amino Acid, i, 1)
Flag = True
Set rs = db.OpenRecordset ("Select pa from Amino Acid where Amino acid=" &
strAnAmino & "")
If rs.RecordCount > 0 Then
ipa_value(j) = rs. Fields (0)
For k = 0 To 3
If IPA value (k) \ge 100 or ipa_value (k) = 0 Then
Flag = False
Exit For
Next k
If flag = True Then
Call BackwardFindAlphaHelicesPartII1 (Alpha partl, i)
Exit For
Else
If j>=3 Then
For k =0 To 2
ipa_value (k) ipa value (k + 1)
Next k
J=2
End if
      End if
I=j+1
      End if
           Next i
End Sub
_____
Flag = False
Exit For
End if
Next k
      If flag= True Then
      IApa part1 From Where
      FindAlphaHelicesPartll (From Where)
      Fork 0 To 5
      ipa_value(k) = 0
Next k
j3=1
Else
If j \ge 5 Then
For k = 0 To 4
ipa_value(k) = ipa_value(k + 1)
Next k
```



J=4 End if End if

J=j+1 J=j+1 From Where = From Where + I Wend End Sub

Private Sub FindAlphalHelicesPartII (Alpha_partl As Long) Dim ipa_value (3), c, i, j, count As Long Dim strAnAmino As String For i= Alpha_part1 +1 to Len (Amino Acid) StrAnAmino = Mid (Amino Acid, i, 1) Flag = True Set rs = db.OpenRecordset ("Select pa from Amino Acid where Amino acid=""&StrAnAmino&"")

If rs.RecordCount>0 Then ipa_value(j) = rs.Fields(0) For k =0 To 3 If ipa_value(k) >= 100 Or ipa_value(k) 0 Then flag = False

Private Sub ForwardFindAlphalHelicesPartIII (Alpha_partl as Long, By Val up to As Long) Dim Sigma_Pa, Sigma_Pb, Region_ Count, c as Long Dim strAnAmino As String Dim rs As Recordset Sigma_Pa = 0 Sigma_Pb = 0 On Error GoTo Solve For i = Alpha_partI +1 to up to STR An Amino = Mid (Amino Acid, i, 1) Set rs = db.OpenRecordset ("Select pa, pb from Amino Acid where Amino acid=" &

strAnAmino & ") Sigma_Pa = Sigma_Pa + rs. Fields (0) Sigma_Pb = Sigma_Pb+ rs. Fields (1)

Next i

Region_Count = up to - Alpha_part1 If Region_Count >5 and Sigma_Pa > Sigma_Pb Then Alpha_Helices = Alpha_Helices + Region_Count Alpha Range (ARange_Count, 1) = up to

Alpha Kange (AKange_Count, T) = End if

Exit Sub

Solve: MsgBox "Error in Forward Alpha"



Private Sub Form _Load0 Me.Left =3800 Me.Top=0 Me.Height=4425 Me.Width 8150 IF i Flag="MAIN" Then MsgBox "Select File and Open save Amino Sequence" MnuAmino. Enabled False Else MsgBox "Select Find Menu and Amino Acid Sequence" End if

End Sub

Private Sub Form _Resize () On Error Resume Next txtshow.Height= Me.ScaleHeight-500 txtshow.Width =Me.Scale Width End Sub

Private Sub mnuAmino Click () PB.Visible True MnuSec.Enabled True MnuBack. Enabled True LOSOURCESTR FrmMrna.txtMrna.Text LOSOURCESTR FindStartCodon (LOSOURCESTR) If LOSOURCESTR "" Then PB.Max Len (LOSOURCESTR) * 2 Filling (LOSOURCESTR) End if End Sub

Private Sub mnuBack_Click () IFlag = "AMINO" get Amino = "" Get amino strAmino FrmPB.Show FrmBackProcess.Show End Sub

Private Sub mnuCopy_Click () Clipboard.SetText (txtshow.Sel Text) End Sub Clipboard.SetText (Ixtshow.SelText)

Private Sub mnuCut ClickO Clipboard.SetText (txtshow.SelText)



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Extshow.SelText End Sub

Private Sub mnuExit Click () Unload Me End Sub _____ _____ Private Sub mnuOpen Click () MnuAmino. Enabled = False MnuBack. Enabled = True mnuSec.Enabled = True Dim LinestrAmino, char Amino as String Dim no, k as Integer On Error Resume Next cd1.ShowOpen txtshow.Text = "" txtshow.LoadFile (cd1.FileName) On Error Resume Next Open cd1.FileName For Input As #1

Do Until EOF (1) Line Input #1, LinestrAmino StrAmino = strAmino & LinestrAmino StrAmino = strAmino & ":" Loop Close #1 End Sub

Private Sub mnuPaste_Click () txtshow.SelStart =Clipboard.GetText () End Sub

Exit For End If Next k If flag =True Then Call ForwardFindAlphalHelicesPartlII (Alpha_part1, i) From Where i Exit For Else If> 3 Then For k 0 To 2 $ipa_value(k) = ipa_value(k + 1)$ Next k j=4 End If End if J=j+1End if From Where=From Where+1 For i=0 To 3



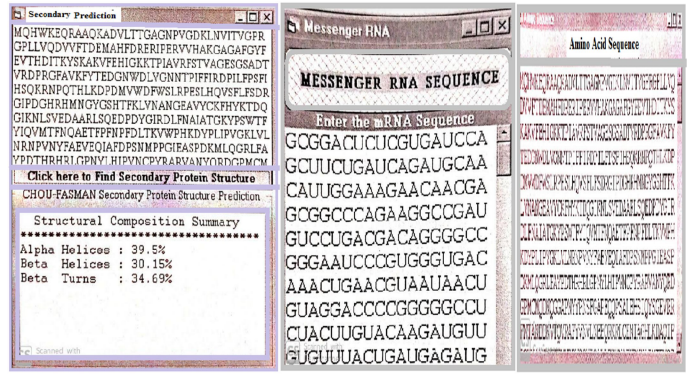
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End ipa value (i) = 0Next i For i = (Alpha partl - 6) To WhereUpTo Step -1 StrAnAmino = Mid (Amino Acid, i, 1) Flag = True Set rs = db.OpenRecordset ("Select pa from Amino Acid where Amino acid=" & strAnAmino & "") If rs.RecordCount > 0 Then $ipa_value(j) = rs.$ Fields (0) For k = 0 To 3 If ipa value $(k) \ge 100$ Or ipa_value (k) = 0 Then Flag = False Exit For Next k J=0If flag = True Then Call BackwardFindAlphaHelicesPartII1 (Alpha partl, i) Exit For

IV. RESULTS AND DISCUSSION.

An amino acid sequence catalase was taken from Swissprot and the E-score value was noted to be negative from BLAST. Then the sequence was saved in visual basic program and noted for the secondary structure prediction of catalase and their back translation. The below result shows the amino acid sequence of catalase, secondary structure prediction alpha, beta, and turn content of catalase., back translation process and sequence conversion was also found using visual basic computer programming.

Recently a new method called the self-optimized prediction method(SOPMA)has been described to improve the success rate in the prediction of the secondary structure of protein. In this paper we report improvemnts brought about by predicting all the sequence of a set of aligned proteins belonging to the same family.





International Journal for Research in Applied Science & Engineering Technology (IJRASET)

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Gibrat (GOR3) result for (Catala	se						SOPMA ret	sult for Catalase	30	40	so	50 ? 1
13 29 MORE PLAND CONTROL AND	ADVILTI bbbsei prission	VFER hhtt NSNP DQGI PLIP NVLIP eth hhttp hhttp	A OTHLES POTHLES A POTHLES A POTHLES A POTHLES A A A A A A A A A A A A A	EDAA HANN EDAA HANN HISTO HIGT HIGT HIGT	NETVARESCEAT NETVARESCEAT NETVELESSEEN NETVELESSEEN NETVELESSEEN NETVELESSEEN NETVELESSEEN NETVELESSEEN NETVELESSEEN	TYSPLFSFAN TYSPLFSFAN LPSALATOR LPSALATOR TSSBCPCI TSSBCPCI TSSBCPCI TSSBCPCI	AND	RVMARIAN booshit weekti weekti ht constitut of dif of dif bit of RLPATPOT bit of SIQYSUEVER	POYFEVTROSTRY CONCEPTION IFPIRDPIL/FGTIN CONCEPTION	Anton 11 HORSNFOT M TTDOGISM CFYFAFVA CFYFAFVA MUNING Abhibhan Mathaba	estence hhttp: http://www.aris. howaris. h	PAGLEREELHO PAGLEREELHO CEDEDEUGIEDL CEDEDEUGIEDL II hhim NFAEVEDIAFD hhimhhh	I LOUIDERATI INTERNATIONAL INO
COR3 :													
Alpha helix	(Hh)				37.83%			30500		(Rb) =	161 18	30.618	
310 100 110	(0g)		0		0.003		_		ipha helix	(09) =	o is		
P1 nerra	(II)			is	0.00%		_		helix helix	(11) =	o is		
Beca Diruge	(SP)			18			_		ta bridge	(8b) =		0.00%	
Extended strand	(Ee)	5	110		20.91%		_	10	tended strand	(Ec) :	81 18		
Beta turn	(TE)	1	0	15	0.00%		_		a turn	(TEJ I	35 20		
Bend region	()	=	0	10	0.00%		_		d region	():	0 is		
Random coil	()	=	217	is	41.25%		_		dom coil	() :		47.34%	
Ambigous states	(?)	1	-	is	0.00%			Anto	igous states	(?) :	o is		
Other states		=	0	is	0.00%			Oth	er states	1	0.14	-	

Secondary Structure rates from Chou Fasman method, GOR and SOPMA

	Chou fasman		GOR Meth	SOPMA	
	method	GOR1	GOR3	GOR4	Method
Alpha helices	39.5%	34.60%q	34.60%	37.83%	29.66%
Beta sheet	30.15%	0.00%	37.83%	20.91%	16.16%
Beta turn	34.69%	13.69%	13.88%	0.00%	7.60%

The results of secondary structure prediction of proteins from chou fasman algorithm using the visual basic method were compared with the GOR and SOPMA methods to show the percentage similarity between the three methods.

Procession in the second se								Back Translation Process For Catalase
GOR4 result for 6	Catala	se						BRNA SEQUENCES
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V. CONCLUSION

The secondary structure of protein catalase was predicted using visual basic programming language. The method of using programming language has been improved and gives the same result like database. Its recognition rate is already equivalent to those of the current best prediction methods. Different colors can also be used to differentiate the model. The regions corresponding to the alpha, beta, and turns can be identified on the basis of sequence length. The individual amino acid corresponding to the secondary structure states can be noted.

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IMPACT FACTOR: 7.129







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