ROLE OF COMPUTATIONAL BIOLOGY IN ORAL SCIENCE

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ABSTRACT

DNA sequence Cigarette Smoking, Betel leaf chewing, and alcohol consumption are major cause of oral cancer in Asia. The difficulty in quitting, coupled with patients' economic conditions affects the inability to get diagnosed early, driving death rate higher. There has been major advancement in molecular sciences, computational biology, and other fields today, but we are not still able to pinpoint the causes of oral cancer, also known as Squamous Cell Carcinoma (OSCC). Early detection leads to better survival rate, therefore, education on yearly check-ups plays a vital role. Computational analysis at the genomic (DNA sequence) can help patients with targeted cellular treatment and hopefully a cure. In this paper, we would look at computation tools used in detecting OSCC and various analysis. Analysis includes detecting abnormality in the cell and other molecular reactions which later morph into a cancerous cell. Later, we investigate all computational tools or techniques from local and global sequence alignment, protein structure, gene, functional structure analysis which help medical staff detect cancer, which in turn can help with oral cancer treatment, prognosis and hopefully a cure.

KEYWORDS

Computational biology, Sequence alignment, Genomic analysis, and Gene

1. INTRODUCTION

Oral squamous cell carcinoma (OSCC) is one of the common types of cancer of head and neck squamous cell carcinoma (HNSCC) today accounting to 500,000 new cases diagnosed and approximately every 250000 deaths every year [1]. Its prevalence is different in different continents or places. the oral cancer is most prevalent in the Asian continent than in the industrialized, more informed western countries where it accounts for 1- 2% of the overall cancer [2]. Oral cancer mortality is highest in Asia unfortunately [3]. OSCC forms almost80 - 90% of all head and neck cancer in the Asian continent [3]. There are many factors which lead to the oral cancer. in the western world, its smoking and alcohol [4].

In Asia, however, the consumption of local liquor, smokeless tobacco and cigarettes with no filters are the leading cause of oral cancer [5]. Gene mutation and the activation of certain genes

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in the cell are seen as early stage of oral cancer but it is not yet proved categorically [6][7]. OSCC is commonly seen in elderly males in the larynx and lips area, accounting to almost 95% of the region [8]. With a 50% survival rate after detection [7], the need to develop newer tools and technologies for early faster detection is greater today [3]. In the next section, we will be talking about the role of computational biology or bioinformatics in diagnosis, and treatment of OSCC.

2. COMPUTATIONAL BIOLOGY

Computational biology or Bioinformatics is a field which encompasses both biology and computer science. This field provides various tools in storing big data sometimes running into petabytes of data stored in databases, and efficiently retrieving the data from the database to establish a relation, provide a 3d structural analysis of proteins and today throws light on the cellular functionality in a much deeper level than what it was prior. This is truer after the genome project was started in the early 1990's which exploded the amount of data stored [9]. The field can also be viewed as a field which brings more analytical power, which in turn would make biological data analysis easier production more biological knowledge [10] [11].

In other words, Computational biology enables scientists to go through huge data easily, finding useful encoded gene segments (for example, a human genome is 3 billion characters in length, a gene could be 100k character in length) [12]. This information is then used in targeted therapy and pharmaceuticals to develop targeted drugs. One of the most pervasive uses of genome sequencing is used to find hereditary relationships and what ethnicity make is one made up of, also known as DNA test / DNA paternity test.

Bioinformatics should not be mistaken to only confined to DNA or genome research augmented study, but it is field which touches a medical science including Molecular Sciences, Drug development, nursing, oral medicine, and other associated fields augmented with pure computer science, biostatistics, and information sciences like State health Care systems or Hospital management systems [9][13].

Computational biology involves developing new data structures, algorithms using either newer and efficient data structures or underpinning hardware coupled with new algorithms to efficiently store new data in the databases, or efficiently retrieve biological relevant data from a huge database. These involve and can vary depending on the algorithm, data mining techniques and visualization or image processing algorithms based off the known biological data from the biologist and lab technicians. Some of the biological niches today are in sequence alignment tools (local and global), multiple sequence alignments, genome development, Gene alignment, protein structure (2D and 3D image development), phylogenetic tree development, and prediction like protein – protein interactions and gene expression. Most of these predictions are based off existing prediction models which are modified based on the known biological knowledge from researchers over the years [14].

3. COMPUTATIONAL BIOLOGY IN ORAL SCIENCE

Oral Science computational biology is a niche field inside Bioinformatics as it is now catering to only head and neck although it still touches upon other molecular sciences like Genome, Gene expression, protein expression, Protein structure and Sequencing methods. Sequencing and Gene sequencing is probably more useful to dentist today than any other bioinformatic tool. Oral science tools will be discussed in detail in the subsequent sections.

Today MEDLINE (Medical Literature Analysis and Retrieval System Online) houses large number of published articles on health. MEDLINE also houses bibliographic information on articles ranging from oral science, general medicine, nursing, general health care, and veterinary science. It is an open source, and most articles can be searched through PubMed. Like MEDLINE [9], there is Latin American one, called The Literature Latino-Americana e do Caribe em Ciências da Saúde (in Portuguese), or LILACS which was called previously called Latin American Index Medicus. There is one other website which is called 'Medscape' which provides health care practitioners including dentists and nurses (other health care professionals) [15]. The primary goal of any oral computational biologists is to improve oral care, which includes better and efficient diagnosis, treatment and prevention of disease and maintain overall good oral health. In regions like Asia, where one needs to see the economical angle, Oral computational biology should also strive to delivery cost effective oral care at the same time [16]. Today, we see that, oral science practitioners and scientists are using new technologies which have come out of this fusion between oral / health science and computer science [17].

Oral science pathology is one of the streams of many including orthodontics, periodontics, and others. Oral Pathology is a specialty mostly dealing with the diagnosis and understanding the causes of diseases related to head and necks, jaws, and face overall. Oral pathology is not limited to above said areas but also includes supporting bone, joints gums, tongue, and surrounding tissues. Knowledge of the above is acquired by learnt methods and disciplines which are basic to oral science practice, through microscopic anatomy, microbiology, and physiology, either through literature or knowledge acquired over a period while practicing dentistry on patients. Oral Pathology uses human science best practices, tools, and computational algorithms to diagnose and treat any diseases or abnormalities in the region [18]. Oral pathology especially in Asian, more so in South Asia, takes into consideration current and pass history coupled with diet, hygiene practice, economic condition, and lifestyle risks like smoking, areca nut chewing, alcohol consumption and recently fad called 'vaping' [9].

The abnormalities seen today are mouth sore that fails to heal or bleeds, to reddish and whitish patches inside the mouth, a lump which is growing quickly and thickening, sore throat leading to difficulty in chewing or swallowing. All the said abnormalities could be seen inside the mouth, face, or neck. Such abnormalities could be diagnosed as oral cancer, cysts, tumour (both benign or malignant), salivary gland abnormalities or cancer (benign or malignant) or simple diseases or lesions. Over the years, large data and literature have been published with varying sites, varieties of above said abnormalities. Today with the advent of projects like the Genome project [9], oral pathology and computational biology are converging ever so closely and working towards automated, efficient, and better way to diagnose pathology at the early stages before they turn life threatening thereby increasing the chances of survival or in other cases, improving the lifestyle of the patient.

4. COMPUTATIONAL TOOLS IN ORAL SCIENCE

In this section, we would look at the computational tools available at the oral professional disposal.

4.1. Oral Cancer Diagnosis

Among head of neck squamous cell carcinoma, oral cancer is most common. It is most prevalent in the males, and we see oral cancer making up to 7% of all cancers in Asia especially in south Asia. The cause for this high rate has been discussed earlier. Oral cancer if detected at an early

stage improves the chances of the patient returning back to normal life quickly perhaps with a permanent cure [19] Unfortunately, when the patient presents themself for an examination, more than likely, the cancer would have spread to other regions and lymph nodes and there is very little a practitioner could do to the patient [20]. Hence yearly check-ups are advised. Sometimes, the lesions found in the mouth are not easily discernible to experienced practitioner leading to false diagnosis. Although some lesions are detected, the similarities in benign and malignant is very high leading to poor and false diagnosis [21].

OralCDx which is a brush biopsy method developed in early 1950s was successful in detecting oral cancer using computer assisted sample analysis. The biopsy collects 3 layers of cell epithelium of the oral mucus for better results. The biopsy sample is stained with Papanicolaou tint [22], it is then scanned and analysed microscopically with the help of computer which contains an image database containing different stages of abnormal cell morphology [23]. This computer program is powerful enough to detect abnormal cells among thousands of normal cells [22]. Once the computer prints the data of its analysis, a cytopathologist interprets this analysed data and classifies the sample to atypical (uncertain diagnosis), positive for oral cancer, normal cells, and incomplete sample [24].

To reduce the false negative finding, many attempts were made to the improve the analysis of the application. However, it was tuned to present day success by incorporating neural network computers initially developed for the missile defence [9]. This improved the application's ability to search for abnormal cell from its database and printed out a detailed data for the cytopathologist to interpret this detailed data in an efficient way there by reducing the chances of overlooking any abnormal cells today.

4.2. Genome Research

Genome is defined as the entire genetic markup of an organism or sum of all genes excluding the part of the DNA which we have still not understood [25]. It can also be looked upon as a blueprint for all cellular activities including the protein structure, the region where the gene is expressed inside each cell. This blueprint contains entire set of instructions for new cell creation, construction, operation, maintenance, and repair in an organism [26]. In 2003, human genome project was initiated with a n objective to obtain and study genes, its structure and interactions amongst the genes [9]. Other genome projects were started much earlier like the Drosophila genome, which is comparatively shorter than the human genome which runs to 3 billion characters in length. All genome projects were initiated with a goal to diagnose and prevent diseases. Genome should not be confused with genetics which is a study of genes in an organism.

With new genomic discoveries, researchers and clinicians are throwing more light into our understanding of oral science. These new discoveries coupled with faster computer which are assembled to a supercomputer are helping us understand the oral and dental diseases like never before. Precision diagnostic tools, procedures, tests with new practical based research are throwing new research vectors providing novel avenues to diagnose, treatment (drug discovery) and improve the oral health.

Head and Neck Squamous Cell Carcinoma (HNSCC) is a disease with complex gene alterations, where some genes are mutated with an additional protein character or removed from the position in the gene (An indent '-' is associated when a protein is missing from its known location in a mutated gene). HNSS Tumors associated with human papillomavirus infection have a genetic profile different from the tumors associated with excessive tobacco usage [27. CCND1 gene is associated with oral Squamous cell carcinoma[32]. Overexpression of this CCND1 is a sure sign of aggressive nature, and early recurrence [32]. Over expression has also indicator of the overall

effectiveness of radio [28,29] Erb B complex is a cell cycle signaling gene and its overexpression is an indicator for OSCC [30].

Human cells contain tumor suppressor gene and one such gene is PTEN, if absent is the indicator to the SCC. Others include the growth factor MET oncogene. High expression of epithelial proactive cell nuclear antigen (PCNA) is an indicator for poor survival rate and prognosis [30]. Other genes like UGT1A7, GSTMI, GSTT1 and CYP1A1 over expression is most seen in the smoking and tobacco users [31]. Ever since the genome projects started in the early 1990s coupled with the human genome project in 2003, huge amount of data is collected in either full genome sequences which in a human can run up to 3 billion characters in length to smaller sequences of few hundred thousand like drosophila.

Walter Goad and his colleagues were the first to assemble a DNA Sequence Database in their lab at Los Alamos National Library (LANL) which later came to be known as GenBank [58] [9]. European equivalent was later developed, and it is housed in Germany, called EMBL (European Molecular Biology Laboratory) [58]. Japanese who were the first pioneers for genome research always had their own database called the DNA Databank of Japan at Mishima. Today, all these three have collaborated and formed a global International Nucleotide Sequence Database which is where every technician would go from Genome, protein, or genes [9].

4.3. Proteomics and Computational Science

Proteomics is a new word which is commonly used today, but it was first used I the Therapeutic field. The world Proteome was coined by Marc Wilkins in 1995 [33]. Ever since then, the word 'omics' is attached to anything related to protein and its study. Proteomics is little different from genomics in that, a genome study is about genetic markup or a blueprint for all genes, proteins, and all cellular activity, while proteomics is a study of protein, its structure, expression, and its interaction [9]. All the three, are complex because they all depend on the organism need, time, and environmental conditions [34]. A human genome contains code to about 26000 - 32000 proteins [35], but the number of proteins expressed inside our body is about 1 million.

Today there are many procedures inside proteomics ranging from one and two -dimensional gel electrophoresis [36]. gel-free screening technique called multidimensional protein identification. methods [37], isotope labelling with amino acids [38], isotope-coded tags, isobaric tags, and their quantitation [39]. There are other methods like shotgun-proteomics [40], gel electrophoresis [41] and microarray for proteins [42], all these techniques or procedures can be used on cells, tissues, and organs [9]. Today, Large-scale western blot assays [43] and multiple reaction monitoring assays [44], are used for high-throughput processing and it is a throwing new light in our understanding of protein interaction, protein synthesis and protein culture.

Tools	Tool Used for		
InterProScan	Used for Protein functional analysis		
PfamScan	FASTA sequence searching against		
	Pfam library		
HMMER3	Used for searching one or more		
	sequence against a sequence database		
Hmmscan	Search other sequences against		
	collection of profiles		
Phobius	Transmembrane topology and signal		
	peptides prediction [58]		
RADAR	Protein sequence alignment		

Table 1: Protein	Structure	Databases	[58]
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Computational biology or algorithms are used in the proteomics domain mainly to manage, store, retrieve markers [45]. These markers could be from DNA sequences or genes or proteins [45]. The algorithms collect and hold massive amount of data sometimes in petabytes and are effective to bridging the gaps between other streams like genomics, metabolism, and protein synthesis. This entire process of providing an association between various niche fields which are closely related (genome, genes, proteins, metabolism, and synthesis) is a computational costly process.

By costly, we mean, computer intensive process, lately proteomics has moved on to heuristics rather than dynamic programming algorithms. We will talk about these in detail in the sequence alignment algorithm section. The algorithms effectiveness in terms of sensitivity and making good biological sense is exacerbated by different parameters, the definition of quality and sensitivity definition and the standard data formats. In other words, there is no standard yet when it comes what makes an algorithm effective and what is the standard data on which algorithms are compared with [46]. A collection of reactions inside the cell that have a biological impact are called protein pathways [47], there are many tools to determine these pathways today [47]. Some of the databases are in place today such are Kyoto Encyclopedia on Genes and Genomes, BioCarta, pathway Reactome and Ingenuity are some of the databases which have knowledge on cell reactions, metabolism, and cell - cell interactions, protein synthesis and protein - protein interaction [48, 49]. PANTHER is another database, which concentrates more on protein analysis based off the evolutionary relationship. This database also works on genomes to extract these relationships [50, 51]. Cancer related pathways and their detection are stored in a database called Netpath [52]. other protein interaction information is stored in BIOGRID and MINT [53, 53, 54, 55]. Most of this information are extracted using sequence format [56, 57], which we will talk later in this paper. Table 1 shows a collection of databases concentrating on structure analysis.

Tools	Description		
PyMOL	Viewer and modelling		
ERRAT	Verification and evaluation		
PROCHECK	A structure validation web server		
Swiss model	Homology modeling of a protein whose structure was unknown		
RAMPAGE	Examine the quality of protein structure		

Table 2: Protein Structure Tools [58]

Protein structure prediction and their classification is used as a tool for predicting OSCC today. To classify and prediction protein structure, well know knowledge base in the form all know protein structures for an organism is developed and stored [45]. Protein data bank (PDB) is one database which houses proteins, DNA and RNAs in a 3D format. Once OSCC is detected, the conforming proteins can then be examined, analysed and either targeted treatment or drug discovery could then be made. Typically, these methods are either NMR Spectroscopy, X-Ray Spectroscopy or Cryoelectronic microscopy [51]. Structural tools are shown in Table 2.

4.4. Micro Arrays

Microarray is a tool which provides a platform to measure the expression of large set of genes simultaneously. It helps in identification of Single-nucleotide polymorphisms (SNPs), Mutations, tumor classification, target genes to suppress tumor, biomarkers, and identification of genes with chemoresistance. All these help in customized drug treatment and drug discovery [9]. A Microarray can be seen as a circuit made up of thousands of microchips, with each chip containing a short, DNA sequences which together would add up to a normal gene and its

variants in human population [9] [59]. These microchips are not as advanced as to create their own memory operations like mutual exclusion or locking [94]

When the Microarray were first developed, there were primarily used for research to identify anomalies in Individuals. As time progressed and enough data collected using manual method, Microarray were used later to find how often a mutation in a particular gene would end up with a oral cancer [59]. Regions where the mutation were seen in a gene were later correlated with oral diseases, Tumor (Benign or Malignant) classification or other oral diseases. As computer became powerful both in terms processing power and memory, large features and large portion of the genome were placed in each individual chip thus enabling many types of genes mutations [3]. Microarrays are also used today to study which gene is turned on or off inside each cell in an organism. They are also used to isolate RNA from the samples for measurement [59].

Microarrays find their usefulness in the labs for clinical testing of diseases such as COVID [59]. As a result of their usefulness in determining the gene mutations, they are used to develop customized drug for patients, as genes determine how organisms can react to certain chemical drugs developed [59]. Microarrays are quickly fading away because of faster, efficient sequencing algorithms which are still costlier than Microarray technology [9].

4.5. Sequence Alignment

A sequence in molecular or genome research could be either a DNA, RNA, or protein sequence [60]. These sequences are made up of amino acid representing characters. A protein sequence is made up of strings of character (A, R, N, D, C, Q, E, G, H, I, L, K, M, F, P, S, T, W, Y, V). Similarly, DNA (A, C, G, T) and RNA (A, C, G, U) [61]. Sequence alignment is a very old paradigm in the microbiology, where the objective is to find regions of similarity between either 2 pair of sequences or multiple sequences. These similar regions can be analysed later, to determine the similar genes, protein structure and other information leading to drug discovery and treatment [60] [62].

Early research depended on manual efforts to align or find regions of similarity between two sequences. Further research classified pairwise sequencing into local and global sequencing. Local sequence alignment is when regions in short burst are aligned together whereas global sequence alignment is interested in finding all regions of similarity in the entire sequence (DNA, RNA, or protein). Two early algorithms developed in 1970s and 1980s using a technique called dynamic programming, they are Smith-Waterman and Needleman (local alignment) and Wunsch algorithms (global alignment) [63] [64]. The drawback of Dynamic Programming algorithms was that they were very slow. To overcome tis shortfall, Heuristic algorithms were developed. Some of the local heuristic-based algorithms are FASTA [65], BLAST [66], Gapped BLAST [67], BLAT [68], BLASTZ [69], and PatternHunter [70]. Similarly, popular heuristic based global sequence alignment algorithms are MUMmer [71], GLASS [72], AVID [73], and LAGAN [74].

To quantify the alignment in heuristic algorithms, a scoring function is used. When characters in each sequence match or mismatches, a score is given to that character alignment. A scoring function was developed for aligned of pairs of characters, then scores of aligned pairs of characters together were added to get the final sequence alignment score [60]. A scoring matric was later developed, which is a n * n matrix in which each corresponding base pair has a score [75]. PAM (Percentage of Acceptable point Mutations per 108 years) series of matrices [76] [77] and BLOSUM (BLOcks SUbstitution Matrix) series of matrices [78] are popular scoring matrices. All heuristic algorithms use three techniques to find the final alignment, they are look-up table, short alignment of length from 8 - 15, called seeds and finally maximum match subsequence. The look up tables are used by FASTA and BLAST family. Seeds and their variety

are used by BLAT, PatternHunter, Glass, AVID [60-63]. Maximal match subsequence which are a new technique was introduced in LAGAN, and MASAA family [61, 62, 63, 92, 93].

BLAST, LAGAN and MASAA family are currently the most effective of the alignment algorithms as they are fast (BLAST) or relatively and sensitive (LAGAN and MASAA family). LAGAN and MASAA variants use different techniques, data structures and stitching techniques to arrive at the final alignment. All the above 3 algorithms are used in our labs to find local and global alignments. Pairwise alignment can be used to find multiple sequence alignment. Initial multiple sequence alignment started with pairwise alignment and progressive moved to other sequences building a guide tree [79] and creating an unwanted pair group with arithmetic mean [80]. Guide tree is used to dictate how multiple sequence alignment would progress. Closely related sequences are aligned first and once there are all done, divergent sequences are then aligned on top of them. This process is called progressive multiple alignment method. ClustalW [81], MAFFT [82], Kalign [83], Probalign [84], MUSCLE [85], DIALIGN [86], PRANK [87], FSA [88], T – Coffee [89, 90], and Probcons [91] are use progressive methods. The other method is iterative method, as the name implies, the algorithm repeatedly applies Dynamic programming on an already sequenced pair to eliminate errors [96, 97].

Pairwise sequence alignment is used to detect chronic diseases like COVID - 9, oral diseases from benign to Malignant [60] and multiple sequence alignment together are used for protein structure prediction, protein-protein interaction, predicting cellular regions where genes would be expressed, Phylogenetics trees (DNA make up, paternity tests). Today artificial intelligence (AI) is also helping sequencing algorithms to predict tumour growth, or mutations in the head and neck regions [95].

5. CONCLUSION

Computational biology has played a pivotal role in our understanding and treatment of oral science. The potential to quickly diagnosis a benign tumour to malignant is not far from way with the way computation biology is progressing today. There are certain fields which may go extinct as time progresses, Microarrays is one of them. Currently the only advantage it has versus sequencing algorithms is the cost. When more commercial sequencing algorithms come into the field, the cost would dramatically come down. Genomics and proteomics will be the future as genes, and their expression (depends on the region, need and environment) is not fully understood. To understand more in detail, we need to convert huge amount of data into knowledge, this means computational ideas, computer hardware and techniques. The future is bright for drug discovery, customized treatment (gene therapy [98]) and a combination of new cellular discovery knowledge and computational prediction model would unlock better results which perhaps might lead to oral cancer / cancer cure.

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