Bioinformatics and Its Scope in Biotechnology

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Abstract: Bioinformatics is the procurement use of computational instruments, stockpiling, and plan, distinguishing proof, filing, investigation, elucidation perception and correspondence approach for extending the utilization of organic, restorative, conduct or wellbeing information in atomic science explore improvement. The examination in biotechnology particularly that including grouping information the executives and medication configuration happened at a rapid rate because of improvement of bioinformatics. Various apparatuses and programming are produced for investigation and elucidation of natural unpredictability. There are number of uses of bioinformatics viz. succession examination and arrangement, atomic displaying, docking, explanation and dynamic reproduction to quicken the biotechnological inquire about. It is normal that numerous future bioinformatics developments are probably going to animate examination of immense organic information. Here, in this correspondence, we have endeavoured to clarify the significance of bioinformatics in different field of biotechnology viz. genomics, proteomics, transcriptomics, cheminformatics, environmental change contemplates, tranquilize revelation and advancement, squander cleanup, bio energy, crop improvement, veterinary sciences, scientific sciences and biodefense.

Index Terms: Bioinformatics, Biotechnology, Software, Proteomics, Proteomics analysis.

I. INTRODUCTION

Bioinformatics is the short form for ‘Biological Informatics’. This is considered to be an amalgam of biological sciences and computer science and now a day, many scientists prefer to use the term, computational biology. This part of science turned out to be progressively well known when human genome venture appeared. Bioinformatics blends science, software engineering and data innovation to frame a solitary control. It covers numerous territories of natural science particularly of current science viz. genomics, transcriptomics, proteomics, genetics, and evolution. A definitive objective of the field is to empower the revelation of new natural bits of knowledge just as to make a worldwide point of view from which bringing together standards in science can be perceived. Bioinformatics is an intriguing subject having contribution of building workmanship just as of science. Bioinformaticians are for the most part occupied with structuring new calculations, programming, creating refreshed databases which all assistance in tackling numerous natural issues. Various bioinformatics apparatuses, programming and databases are accessible for better comprehension of natural intricacy and investigate and store the organic information. Hence, the bioinformatics explore is utilized to maintain a strategic distance from time, cost and wet lab practice. Researchers understood the significance of arrangement databases in 1950s and that is the reason, first protein grouping database was made in 1956 soon after insulin peptides successions ended up accessible.

The human genome succession information is huge to the point that whenever accumulated in books, the information would keep running into 200 volumes of 1000 pages each and perusing alone would require 26 years working nonstop. This test of dealing with such gigantic information must be conceivable in view of bioinformatics. The development of the biotechnology business lately is exceptional, and progressions in atomic demonstrating, sickness portrayal, pharmaceutical revelation, clinical human services, criminology, and farming generally sway monetary and social issues around the world. Subsequently, with individual’s certainty and advancement of biotechnology, bioinformatics additionally came to higher than ever among all the organic sciences. There exists various uses of bioinformatics for quickening research in the territory of biotechnology that incorporate programmed genome sequencing, quality recognizable proof, expectation of quality capacity, forecast of protein structure, phylogeny, medicate planning and improvement, ID of creatures, antibody structuring, understanding the quality and genome intricacy, understanding protein structure, usefulness and collapsing. By utilizing bioinformatics in research, many long haul ventures are turned up so quick like genome mapping of human and different life forms. So also, it is normal that bioinformatics advancement in future will likewise satisfy the needs of biotechnology.
A. Goals of Bioinformatics

The advancement of proficient calculations for estimating arrangement closeness is a critical objective of bioinformatics. The Needleman-Wunsch calculation, which depends on powerful programming, ensures finding the ideal arrangement of sets of groupings. This calculation basically partitions an expansive issue (the full succession) into a progression of littler issues (short arrangement portions) and uses the arrangements of the littler issues to build an answer for the substantial issue. Similitude’s in groupings are scored in a framework, and the calculation takes into account the recognition of holes in arrangement. In spite of the fact that the Needleman-Wunsch calculation is powerful, it is unreasonably moderate for testing a vast succession database. Along these lines, much consideration has been given to discovering quick data recovery calculations that can manage the immense measures of information in the chronicles. A precedent is the program BLAST (Basic Local Alignment Search Tool). A development of BLAST, known as position-specific iterated- (or PSI-) BLAST, makes use of patterns of conservation in related sequences and combines the high speed of BLAST with very high sensitivity to find related sequences. Another objective of bioinformatics is the expansion of trial information by forecasts. A crucial objective of computational science is the expectation of protein structure from an amino corrosive succession. The unconstrained collapsing of proteins demonstrates this should be conceivable. Advance in the improvement of strategies to anticipate protein collapsing is estimated by biennial Critical Assessment of Structure Prediction (CASP) programs, which include daze trial of structure forecast techniques. Bioinformatics is likewise used to anticipate communications between proteins, given individual structures of the accomplices. This is known as the “docking problem.” Protein-protein complexes show good complementarily in surface shape and polarity and are stabilized largely by weak interactions, such as burial of hydrophobic surface, hydrogen bonds, and van der Waals forces. PC programs mimic these collaborations to foresee the ideal spatial connection between restricting accomplices. A specific test, one that could have imperative helpful applications, is to structure a counter acting agent that ties with high fondness to an objective protein. At first, much bioinformatics investigate has had a moderately limited center, focusing on contriving calculations for dissecting specific kinds of information, for example, quality groupings or protein structures.

II. BIOINFORMATICS TOOLS AND SOFTWARE

1) Blast: Basic local alignment search tool, provided by NCBI.
2) Fasta: This server is hosted by the University of Virginia, USA. It harbors multiple online software for sequence comparison, local and global alignment.
3) Auto Prime: Auto prime is very useful software for designing Reverse Transcription Real Time PCR (Q-RT-PCR) primers that are specific to the exon-intron boundaries.
4) Primo Pro 3.4: A java enabled online primer designing tool.
5) Primo Pro 3.2: It’s another online primer designing software. Its notable feature is that it can reduce background noise by exercising check on mispriming on non-target DNA sequence. It likewise “presents a cluster mode alternative for high throughput PCR groundwork plan”.
6) Meth Primer: Its very useful site for designing primers for methylation PCR (Denatured, single stranded DNA (ssDNA) is modified with sodium bisulfate " pursued by PCR intensification utilizing two sets of preliminaries, with one sets explicit for methylated DNA, the other unmethylated DNA”).
7) Gene Fisher: "Gene Fisher is an interactive web-based program for designing degenerate primers". The basic suspicion is "supposition that qualities with related capacity from various creatures show high succession similitude, degenerate preliminaries can be structured from arrangements of homologues qualities". This suspicion "prompts segregation of qualities in an objective living being utilizing various arrangements of related qualities from various creatures".
8) Codehop: "The Consensus Degenerate Hybrid Oligonucleotide Primers (CODEHOP) program is hosted by the Fred Hutchinson Cancer Research Center in Seattle, Washington and designs PCR (Polymerase Chain Reaction) primers from protein multiple-sequence alignments"
9) Primo Degenerate 3.4: "Primo Degenerate 3.4 designs PCR primers based on a single peptide sequence or multiple alignments of proteins or nucleotides. For savage preliminaries, the likelihood of official to the objective is corresponding to the powerful grouping of the particular preliminary".
10) **Oligo.Net**: "OLIGO Primer Analysis Software is the basic instrument for planning and dissecting sequencing and PCR groundworks, manufactured qualities, and different sorts of tests including siRNA and sub-atomic signals. In view of the most exceptional closest neighbor thermodynamic information, Oligo's scan calculations find ideal introductions for PCR, including, TaqMan exceedingly multiplexed agreement or ruffian groundworks. Different record clump handling is conceivable. It is likewise a significant device for site coordinated mutagenesis".

11) **Array Designer 2**: It's an Oligo and cDNA Microarray Design Software. "It plans tests for SNP recognition, microarray quality articulation and quality articulation profiling. Furthermore, thorough help for tiling clusters and resequencing exhibits is accessible".

12) **Primer Premier**: Primer Premier is one of the "most comprehensive software to design and analyze PCR primers". Primers can be designed for standard PCR, SNP genotyping assays, Multiplexing assays, along with checking the secondary structures of the designed primers.

13) **Gene Copoeia**: "GeneCopoeia, Inc. is a US-based maker and supplier of genomics and proteomics items and administrations for scholastic and legislative research organizations, pharmaceutical and biotechnology industry". "GeneCopoeia offers far reaching instruments for microRNA (miRNA) practical examination so scientists can distinguish, express, approve or knockdown microRNA of intrigue unquestionably. All known human, mouse and rodent microRNA in miRBase secured".

14) **Dot plot (+) - Dot-plot (+)**: software is used to identify the overlapping portions of two sequences and to identify the repeats and inverted repeats of a particular sequence.

### III. APPLICATION OF BIOINFORMATICS IN VARIOUS FIELDS
Bioinformatics is the utilization of IT in biotechnology for the information stockpiling, information warehousing and breaking down the DNA arrangements. In Bioinformatics learning of numerous branches are required like science, arithmetic, software engineering, laws of material science and science, and obviously solid information of IT to break down biotech information. Bioinformatics isn't restricted to the registering information, yet as a general rule it tends to be utilized to take care of numerous natural issues and discover how living things functions. It is the comprehensive application of mathematics (e.g., probability and statistics), science (e.g., biochemistry), and a core set of problem-solving methods (e.g., computer algorithms) to the understanding of living systems.

Bioinformatics is being used in following fields:

1) Molecular medicine
2) Personalized medicine
3) Preventative medicine
4) Gene therapy
5) Drug development
6) Microbial genome applications
7) Waste cleanup
8) Climate change Studies
9) Alternative energy sources
10) Biotechnology
11) Antibiotic resistance
12) Forensic analysis of microbes
13) Bio-weapon creation
14) Evolutionary studies
15) Crop improvement

**A. Biomedicine**
The human genome will affect the fields of biomedical research and clinical prescription. Pretty much every illness has a hereditary part. The finish of the human genome implies that we can scan for the qualities straightforwardly connected with various sicknesses and start to comprehend the sub-atomic premise of these ailments all the more obviously. This new information of the sub-atomic components of malady will empower better medicines, fixes and even precaution tests to be created.
B. **Drug discovery**
Utilizing computational devices to recognize and approve new medication targets, increasingly explicit prescriptions that follow up on the reason not only the indications of the infection can be created. These exceedingly explicit medications will have fewer symptoms than a significant number of the present meds.

C. **Personalized medicine**
Clinical prescription will turn out to be increasingly customized with the advancement of the field of pharmacogenomics. This is the investigation of how a person's hereditary legacy influences the body's reaction to drugs. Today, specialists need to utilize experimentation to locate the best medication to regard a specific patient as those with the equivalent clinical side effects can demonstrate a wide scope of reactions to a similar treatment. Later on, specialists will almost certainly investigate a patient's hereditary profile and recommend the best accessible medication treatment and dose from the earliest starting point.

D. **Preventative Medicine**
With the particular subtleties of the hereditary systems of ailments being disentangled, the advancement of analytic tests to gauge an individual's helplessness to various infections may turn into an unmistakable reality. Preventive activities, for example, change of way of life or having treatment at the most punctual conceivable stages when they are bound to be fruitful, could result in gigantic advances in our battle to vanquish malady.

E. **Gene therapy**
Not long from now, the potential for utilizing qualities themselves to treat ailment may turn into a reality. Quality treatment is the methodology used to treat, fix or even forestall ailment by changing the outflow of an individual's qualities.

F. **Microbiology**
By contemplating microorganisms genome researchers can start to comprehend these organisms at an exceptionally major dimension and disconnect the qualities that give them their remarkable capacities to make due under outrageous conditions. The entry of the total genome arrangements and their capability to give a more noteworthy understanding into the microbial world and its abilities could have wide and expansive ramifications for condition, wellbeing, vitality and modern applications. Biotechnology

1) Waste cleanup
2) Climate change
3) Alternative energy sources
4) Antibiotic resistance

G. **Agriculture**
The sequencing of the genomes of plants and creatures ought to have huge advantages for the agrarian network. Bioinformatics devices can be utilized to look for the qualities inside these genomes and to explain their capacities. This particular hereditary learning could then be utilized to deliver more grounded, drier spell, malady and creepy crawly safe yields and improve the nature of animals making them more beneficial, more sickness safe and increasingly gainful.

1) Crops
2) Insect resistance
3) improving nutritional quality

IV. **BIOINFORMATICS APPLICATIONS IN LIFE SCIENCES AND BIOTECHNOLOGIES**
Life sciences specialists gather and break down a high measure of various sorts of logical information, including DNA, RNA, and amino corrosive groupings, in situ and microarray quality articulation information, protein structures and organic pathways, and natural flags and pictures of assorted starting point. As of late, an abundance of bioinformatics applications in the fields of essential and connected life sciences has changed the worldview of both research and misuse of information. The advancement of novel and ground-breaking bioinformatics apparatuses committed to organic information securing, information mining, and investigation engaged both the essential and connected life sciences inquire about. These bioinformatics advancements length from instruments for
genome comment and capacity forecast, quality articulation examinations, and databases of natural data, to the rising fields of biomedical utilizations of research, including the improvement of new bioinformatics-based gadgets and prescient applications. This exceptional issue is made out of five unique research papers chose after inside and out friend audit. Chosen papers portray novel bioinformatics instruments as well as databases for crucial or potentially connected research in the wide scope of natural and biomedical sciences. Understanding the hereditary control of complex unique attributes is of crucial significance to horticultural, developmental, and biomedical hereditary research.

A measurable mapping structure, called practical mapping, has been created and widely used to describe the quantitative attribute loci (QTLs) or nucleotides (QTNs) that underlie a mind boggling dynamic quality. Notwithstanding, this instrument isn't appropriate when the bends are intricate, particularly on account of no monotonic bends. Along these lines, to defeat this issue, in their work J. Qi and associates propose the earliness list (E-file) to in total measure the earliness degree to which a variable (or dynamic characteristic) increments or diminishes its esteem.

The creators appear by both hypothetical verifications and reproduction ponders that E-list is broader than practical mapping and can be connected to any mind boggling dynamic quality, even those with no monotonic bends. RNA-Seq tests are these days broadly utilized in a wide scope of studies, spreading over from genome-wide quality articulation and administrative systems basic essential physiological characteristics to human pathologies, including malignant growth. Be that as it may, RNA-Seq information investigations are mind boggling and require the utilization of a few unique devices to control and process the recovered information. The work exhibited by F. Russo and partners demonstrates ongoing progressions and curiosities presented in RNASeqGUI, a graphical interface that enables the client to deal with and investigate huge informational collections (gathered from RNA-Seq tests) in a quick, proficient, and reproducible way.

The here introduced form of RNASeqGUI consolidates graphical interfaces with apparatuses for reproducible research, for example, proficient measurable programming, comprehensible report, parallel executions, reserving, and intuitive and web-exploitable tables of results. The Cancer Genome Atlas (TCGA) information entryway is a stage containing tumor quality articulation information, together with clinical data, empowering scientists to accumulate data on critical genomic changes that happen amid the advancement and metastasis of a tumor.

To assist biomedical specialists with identifying quality articulation designs identified with bosom malignant growth survival, H. Zhenzhen et al. built up an online TCGA information examination stage called TCGA4U, giving a perception answer for the investigation of the connections of genomic changes with the accessible clinical information. The creators trust that the utilization of TCGA4U will move increasingly biomedical scientists to investigate the natural components of those qualities and all the more accurately clarify their job in bosom malignancy improvement, preparing for the find of more focused on treatments and help more bosom disease patients. Foreseeing blood-mind hindrance (BBB) pervasion is fundamental for medication plan of particles that demonstration in the focal sensory system (CNS).

Then again, incidentally acting medications must show constrained capacity to cross the BBB and along these lines be without activity in the CNS. Be that as it may, understanding the procedure of penetration is confounded, since mixes can cross the BBB by latent dispersion and additionally dynamic transport. As an option in contrast to obtrusive creature tests, in silico screening strategies have been acquainted with aid the advancement of focal sensory system dynamic medications. In their paper, D. Zhang and partners portray the plan and execution of a hereditary calculation to foresee the BBB saturation capacity of a given particle, accomplishing more precise outcomes than as of now accessible models. A solitary nucleotide polymorphism (SNP) is the after-effect of the variety of a solitary nucleotide at a particular position in the genome. Other than presenting some level of hereditary variety inside a populace, certain SNPs have been related with the powerless of the person to explicit maladies. The aggregated learning coming about because of the accessibility of human genome successions and the relationship of explicit SNPs with specific maladies provoked the improvement of the alleged prescient preventive customized drug. In their article, P. Ponomarenko and associates examinations the impacts of SNPs happening at the advertiser locale of explicit qualities on the partiality of the TATA-restricting proteins to the advertiser area utilizing their web-administration SNP_TATA_Comparator. All through their work, the creators give a few models and examine how to utilize the bioinformatics application SNP_TATA_Comparator to dissect and separate commented on SNPs from the database “1000 genomes.” Seventeen epic hopeful SNP markers, putatively connected with a few illnesses, are accounted for.
A. Genomics
The investigation of qualities and their demeanour is called as Genomics. This field produces a huge measure of information from quality arrangements, their interrelation and capacities. To deal with this tremendous colossal information, bioinformatics assumes a vital job. With the total genome successions for an expanding number of life forms, bioinformatics is starting to give both theoretical bases and handy strategies for identifying foundational utilitarian practices of the cell and the living being. Bioinformatics assumes a crucial job in the territories of auxiliary genomics, utilitarian genomics and wholesome genomics.

B. Proteomics
The investigation of protein structure, capacity, and collaborations created by a specific cell, tissue, or life form is called as proteomics. It manages systems of hereditary qualities, organic chemistry and sub-atomic science. Propelled methods in science prompted amass tremendous information of protein-protein connections, protein profiles, and protein action example and organelles structures. This huge information can be overseen and access effectively by utilizing bioinformatics devices, programming and databases. Till now, numerous calculations in the field of proteomics viz. picture examination of 2D gels, peptide mass fingerprinting and peptide discontinuity fingerprinting have been created.

C. Transcriptomics
The investigation of sets of all dispatcher RNA particles in the phone is called as transcriptomics. This can likewise be called as Expression Profiling where DNA microarray is utilized to decide the articulation dimension of mRNA in a given cell populace. The microarray system produces immense measure of information, single run creates a large number of information esteem and one investigation requires several runs. Investigation of such tremendous information is finished by various programming bundles. Along these lines, bioinformatics is utilized for transcriptome examination where mRNA articulation levels can be resolved. RNA sequencing (RNAseq) likewise has been incorporated under transcriptomics. It is completed utilizing cutting edge sequencing to decide the nearness and amount of RNA in an example at a given time. It is utilized to dissect the ceaselessly changing cell transcriptome.

D. Cheminformatics
Cheminformatics (substance informatics) centers around putting away, ordering, seeking, recovering, and applying data about concoction mixes. It includes association of concoction information in a coherent structure to encourage the recovery of compound properties, structures and their connections. Utilizing bioinformatics, it is conceivable through PC calculation to distinguish and fundamentally adjust a characteristic item, to structure a compound with the ideal properties and to evaluate its helpful impacts, hypothetically. Cheminformatics investigation incorporates examinations, for example, closeness seeking, bunching, QSAR displaying, virtual screening, and so forth.

E. Evolutionary Studies/Phylogenetics
The investigation of transformative relationship among people or gathering of life forms is characterized as phylogenetics. Taxonomists locate the transformative relationship utilizing different anatomical techniques that takes a lot of time. Utilizing Bioinformatics, phylogenetic trees are developed dependent on the grouping arrangement utilizing different techniques. Different algorithmic strategies are created for the development of phylogenetic tree that are utilized relying upon the different transformative genealogies.

F. Veterinary Science
Nourishment generation from animals can satisfy need of human populace for sustenance. For better bio-economy, there is a need of proficient creature generation and propagation. This is accomplished with better comprehension of animal’s species. Present and new techniques in domesticated animals species utilizing information from trial or field considers with bioinformatics are helping in understanding the frameworks hereditary qualities of complex attributes and give organically significant and precise forecast. At long last, practically the majority of the following age’s omics devices and strategies that are utilized in different fields of organic sciences can likewise be utilized in veterinary sciences.
G. Forensic Science

Measurable science incorporates the examination with respect to recognizable proof and relatedness of people. It is characteristically interdisciplinary with bioinformatics as both are subject to software engineering and measurements. This field depends on the atomic information and numerous databases are being created to store the DNA profiles of known guilty parties. This field is being pushed because of innovative and factual advances in microarray, Bayesian systems, AI calculations, TFT biosensors and others. This gives the compelling method for proof association and derivation.

H. Biodefense

Biodefense incorporates measures to re-establish bio security to a gathering of living beings that are exposed to organic dangers or irresistible infections (in setting of bio-war or bioterrorism). Today, bioinformatics limitedly affects measurable and knowledge activities. There is a need of more calculations in bioinformatics for biodefense so the created databases may demonstrate interoperability with one another. So as to use cutting edge genome sequencing for legal task, bio-risk mindfulness, moderation and restorative insight, there is a requirement for advancement of increasingly computational applications.

V. CONCLUSION

Bioinformatics has got significant effects on biotechnology and its applications. The huge measure of information created by human genome venture or by other genome sequencing task would be unmanageable without the bioinformatics procedure. Without bioinformatics taking care of, translation of this information would be incomprehensible. Due to the bioinformatics application cost of all these real tasks descends definitely.

Bioinformatics also quickened the drug discovery, vaccine design and also the design of anti-microbial agents. Bioinformatics is additionally used to comprehend quality and furthermore genome. Bioinformatics developers are additionally used to look at quality pair arrangement, which helps in recognizing elements of quality and furthermore genome usefulness.

REFERENCES

[1] Shinozaki K, Yamaguchi SK, Seki M, et al. Regulatory network of gene expression in the drought and cold stress responses.CurrOpinPlant Biol. 2003;6(5):410-7.
[2] Bartels D, Sunkars R. Drought and Salt Tolerance in Plants. CritRevPlant Sci. 2005;24(1):23-58.
[3] Stewart, H. 2005. Biomolecular Interaction Network Database. Bioinformatics 6, 194-198.
[4] Bornberg-Bauer, E., and Paton, N.W. 2002. Conceptual Data Modelling for Bioinformatics. Bioinformatics 3, 166-180.
[5] Wong, L. 2002. Technologies for Integrating Biological Data. Bioinformatics 6, 389-404.
[6] Schonbach, C., Kowalski-Saunders, P., and Brusic, V. 2000. Data Warehousing in Molecular Biology. Bioinformatics. 1, 190-198.
[7] Clark, T., Martin, S., and Liefeld, T. 2004. Globally Distributed Object Identification for Biological Knowledgebases. Bioinformatics 5, 59-70.
[8] Lee, T. et. al. 2006. BioWarehouse: a Bioinformatics Database Warehouse Toolkit. BMC Bioinformatics 7, 170.
[9] Bergeron, B. 2002. Bioinformatics Computing. 80-101.
[10] Gautham, N. 2006. Bioinformatics: Databases and Algorithms. 26-53.
[11] H. O. Habashy, D. G. Powe, E. Glaab, N. Krasnogor, J. M. Garibaldi, E. A. Rakha, G. Ball, A. R Green, C. Caldas, I. O. Ellis, RERG (Ras-related and oestrogen-regulated growth-inhibitor) expression in breast cancer: A marker of ER-positive luminal-like subtype, Breast Cancer Research and Treatment, 128(2):315-326, 2011.
[12] E. Glaab, J. M. Garibaldi and N. Krasnogor. ArrayMining: a modular web-application for microarray analysis combining ensemble and consensus methods with cross-study normalization, BMC Bioinformatics,10:358, 2009.
[13] International Human Genome Sequencing Consortium. Initial sequencing and analysis of the human genome. Nature. 2001;409:860–921.
[14] Fraser CM, Gocayne JD, White O, Adams MD, Clayton RA, Fleishmann RD, et al. The minimal gene complement of Mycoplasma genitalium. Science. 1995;270:397–403.
[15] The C. elegans Sequencing Consortium. Genome sequence of the nematode C. elegans: a platform for investigating biology. Science. 1998;282:2012–2018.
[16] Myers EW, Sutton GG, Delcher AL, Dew IM, Fasulo DP, Flanagan MJ, et al. A whole-genome assembly of Drosophila. Science. 2000;287:2196–2204.
[17] Stein L. Genome annotation: from sequence to biology. Nat Rev Genet. 2001;2:493–503.
[18] Benton D. Bioinformatics—principles and potential of a new multidisciplinary tool. Trends Biotech. 1996;14:261–312.
[19] Maggio ET, Ramnaranay K. Recent developments in computational proteomics. Trends Biotech. 2001;19:266–272.
[20] Burley SK, Almo SC, Bonanno JB, Capel M, Chance MR, Gaasterland T, et al. Structural genomics: beyond the human genome project. Nat Genet. 1999;23:151–157.
[21] Tsoka S, Ouzounis CA. Recent developments and future directions in computational genomics. FEBS Lett. 2000;480:42–48.
[22] Debouc C, Metcalf B. The impact of genomics on drug discovery. Annu Rev Pharmacol Toxicol. 2000;40:193–208.
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