Rising Strengths Hong Kong SAR in Bioinformatics

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Abstract Hong Kong’s bioinformatics sector is attaining new heights in combination with its economic boom and the predominance of the working-age group in its population. Factors such as a knowledge-based and free-market economy have contributed towards a prominent position on the world map of bioinformatics. In this review, we have considered the educational measures, landmark research activities and the achievements of bioinformatics companies and the role of the Hong Kong government in the establishment of bioinformatics as strength. However, several hurdles remain. New government policies will assist computational biologists to overcome these hurdles and further raise the profile of the field. There is a high expectation that bioinformatics in Hong Kong will be a promising area for the next generation.

Keywords Bioinformatics research · Bioinformatics education · Government role · Hong Kong SAR

1 Introduction

Concerning Hong Kong’s economic development, Tony Fu-Lai Yu, an economics scholar from the University of New South Wales, Australia, once wrote: ‘From barren island, Hong Kong changed into the mart of East Asia’. But what puzzled economists is that Hong Kong is only a small region with approximately 6 million people living in an area of around 1064 km² [1]. Today, the population of this city has grown to more than 7 million [2]. Hong Kong, a special administrative region (SAR) of the People’s Republic of China (PRC), is situated in the southern part of China that is enclosed by the Pearl River Delta and the South China Sea [http://en.wikipedia.org/wiki/Pearl_River_DeltaPearl River Delta and the South China Sea (http://www.censtatd.gov.hk/FileManager/EN/Content_810/geog.pdf)]. Hong Kong is one of the world’s leading financial and trading centres and is also well known as a free trade zone with low taxation.

The area became controlled by the Chinese in 221 BC, during the Qin dynasty of China [3], and remained under Chinese control until after the Anglo-Chinese War of 1839–1842 (also known as the First Opium War) when it became a protectorate of the UK. The territory of Hong Kong was initially restricted to Hong Kong Island, with the British then extending their boundaries in two phases. The first phase, in 1860, saw the British outpost being extended to the Kowloon Peninsula, and in 1898 it was again extended to include the New Territories. Hong Kong underwent its first peaceful and democratic transfer of power in 1997 when China regained sovereignty [4, 5]. The area had been occupied by the Japanese in 1941 after the Battle of Hong Kong, which occupancy ended after 3 years and 8 months [6] when Japan surrendered at the end of the Second World War, in 1945.
Hong Kong has now become a major global financial centre and one of East Asia’s ‘Economic Tigers’. The territory is economically and technically in an excellent position to assist developing nations worldwide [7]. During the past two decades, the Hong Kong SAR government has been actively pursuing growth in high-tech industries such as telecommunications, biotechnology, information technology and electronics to help develop this economic leader of Asia into a global technological giant in today’s high-tech world.

In Hong Kong, computational biology has adopted bioinformatics as a research and development (R&D) area in its own right. Researchers in the life sciences, biochemistry, molecular biology and computer science have all used bioinformatics as a tool to solve their particular research problems. In this paper, we discuss how the economic growth of Hong Kong and its population age structure have helped to establish computational biology in the territory and have helped with the location of its foundations. To this end, we identify the learning activities, landmark research, activities of bioinformatics companies and role of the government in Hong Kong in establishing computational biology as strength. We have tried to be comprehensive and apologise to any researcher or research group that has not been included in this report.

2 Hong Kong’s Economic Boom and Population Age Structure Assist the Establishment of Computational Biology

Hong Kong has a free-market economy which is highly dependent on international trade and finance. The USA is Hong Kong’s second-largest export partner after mainland China, which is an important factor in the Hong Kong economy (http://www.tid.gov.hk/english/trade_relations/mainland/trade.html). During the past decade, the GDP of Hong Kong has grown steadily and is now about 6.0% (Fig. 1). The territory has one of the highest per capita incomes in the world. This dynamic economic environment provides an added advantage in high-tech R&D areas such as computer science and computational biology [8, 9].

The economy of any nation depends on its age structure, which presents various challenges. The age structure of a population is directly related to that society’s productivity and economy [10]. Hong Kong has an advantage in that the percentage of its population that is of working age is one of the highest in the world. Hong Kong’s working-age population (15–64 years) is about 74.8%, and the median age in the territory is 43.4 years. The older age group (65 years and over) comprises about 13.5% of Hong Kong’s population (Fig. 1). It has been observed that a higher proportion of older people leads to a critical shortage in manpower and tends to undermine productivity. The healthcare costs of an older population are noted to be economically unproductive, which is also a burden for society [10].

Factors such as these highly competitive business environments, the knowledge-based economy, the best-performing economy in Asia in terms of investment and the vibrancy of the working-age group assist in Hong Kong’s rapid growth to become a strong economic force in East Asia and to emerge as a global technological giant able to emphasise its pioneering success in computational biology.

3 Foundations of Bioinformatics in Hong Kong

The initial step in establishing bioinformatics was the commencement of the Hong Kong government’s innovation and technology development programme in 1998–1999, which provided funding of about $5 billion through the Innovation and Technology Fund (ITF). Four types of project are financed through the ITF programmes of innovation and technology support, university–industry collaboration, general support and small entrepreneur research assistance, all involving innovation or technology that should modernise the industry (http://www.gov.hk/en/about/abouthk/factsheets/docs/technology.pdf).

In the late 1980s, especially 1988, work on bioinformatics in Hong Kong was begun by a group of researchers in the life sciences. Two projects are of particular note: first, the use of microcomputers in histopathology [11] and second, the development of the commercial Microbact 24E (MB24E) microsystem for the identification of common clinical isolates of Enterobacteriaceae and non-fermenting Gram-negative bacilli [12]. In 1998, two more bioinformatics papers by different research groups were published in the journal Bioinformatics. Smith and Xue, from the Biochemistry Department of the Hong Kong University of Science and Technology (HKUST), developed a method for summing up and presenting the information contained in a set of aligned sequences to identify patterns within the sequences and represent them in a more accurate and graphical form [13]. Chau et al. [14], from the Department of Applied Biology and Chemical Technology at the Hong Kong Polytechnic University (PolyU), developed a software package entitled ‘TLCQA’ for low-cost analysis of thin-layer chromatography images. Computational biology had thus already spread to the different disciplines within Hong Kong.

The Hong Kong Bioinformatics Centre, jointly established by the Chinese University of Hong Kong (CUHK) and the HKUST in 1998 and funded by the Innovation and Technology Commission (ITC) of Hong Kong, was the first
bioinformatics research centre in the Hong Kong SAR. This research centre acts as a central source of information to support the biotechnology community in this region; its current director is Professor Tsui Kwok Wing Stephen (http://www.hkbic.cuhk.edu.hk/).

4 Bioinformatics Education, Workshops and Conferences in Hong Kong

Universities in the Hong Kong SAR, such as the University of Hong Kong (HKU) (www.hku.hk), the CUHK (www.cuhk.edu.hk), the HKUST (www.ust.hk), the Hong Kong Baptist University (HKBU) (www.hkbu.edu.hk), the City University of Hong Kong (CityU) (http://www.cityu.edu.hk/) and the PolyU (www.polyu.edu.hk), assist in providing manpower training and development in the field of computational biology (Fig. 2). Currently, several undergraduate or postgraduate programmes or courses are available in the specific area of bioinformatics (Table 1). The departments of Biochemistry and Computer Science at HKU offered the first undergraduate programme, a B.Sc. in Bioinformatics, in September 2001; this combined programme of biochemistry and computer science was supported by both departments. This course is not currently being offered by HKU. Instead, HKU’s Department of Biochemistry offers ‘Sequence Bioinformatics’ (course code: BIOC3608), a course for B.Sc. students that carries 6 credit points. This course was planned to enable students to gain a full knowledge of sequence bioinformatics. Outcomes of the course include knowledge of algorithms for sequence alignments, BLAST searches and construction of phylogenetic trees. A course with the same name but a different course code (BIOC3605) is offered by the Faculty of Medicine at HKU for a Bachelor Degree in Medical Science. Although this course also carries 6 credit points, the course syllabus is different; students learn about computational methods for analysis of DNA and protein sequences. In addition, the Computer Science Department in the Faculty of Engineering at HKU offers a course entitled ‘Computational Molecular Biology’ (course code CSIS3026), also carrying 6 credit points, for students pursuing a Bachelor of Engineering degree. The course teaches the students how to use algorithms to work out computational problems related to the enormous...
amount of information produced by modern molecular biology techniques.

In September 2011, the CUHK began to offer a Master of Science (M.Sc.) programme and a postgraduate diploma (PGD) course in genomics and bioinformatics, as part-time programmes taught by the division of Genomics and Bioinformatics within the School of Biomedical Sciences. The M.Sc. programme is a two-year course carrying 24 credits while the PGD course is for 1 year with 15 credits. In both programmes, students learn fundamental subjects such as an introduction to programming, an introduction to molecular biology and genetics, biocomputing and theories and algorithms in bioinformatics. The M.Sc. programme includes some additional mandatory courses such as systems biology and genome informatics, and a research project carrying 3 credits. Two other courses related to computational biology are available in the CUHK’s undergraduate programme leading to a Bachelor of Engineering degree in computer science: ‘Algorithms for Bioinformatics’ (course code: CSCI3220) and ‘Topics in Bioinformatics and Computational Biology’ (course code: CSCI5050). Course CSCI3220 includes topics related to sequence alignment and multiple sequence alignment using different approaches such as Markov property, recursive functions, dynamic programming, FASTA and BLAST and typical clustering algorithms for microarray analysis. Course CSCI5050 includes the topics of molecular biology, data mining, data processing, sequence alignment and biological networks.

The HKUST does not have a separate programme for computational biology. However, there is one course available in the postgraduate programme of the Department of Electronic and Computer Engineering, ‘Introduction to Bioinformatics Algorithms’ (course code: ELEC 5810), an introductory course on computational biology at the molecular level.
The CityU, founded in 1984 as the City Polytechnic of Hong Kong, has been a fully accredited university since 1994. The Department of Computer Science offers a course entitled ‘Computational Biology and Bioinformatics’ (course code: CS4465), which aims to introduce students to the concepts and techniques used in computational biology to develop the practical skills required to solve problems in this domain.

Several workshops have been arranged in Hong Kong for training of manpower in the area of computational biology. Examples of such workshops include: (1) ‘Principles and Applications of Bioinformatics’, held on 22–23 September 1998 and organised by the Department of Biochemistry of the HKU; (2) ‘Phylogenetics and Bioinformatics’, held on 1–4 April 2008 in the School of Public Health at the CUHK; and (3) ‘Bioinformatics Algorithms and Core Technology’, held on 5 September 2012 and arranged by the Department of Computer Science and the Centre for Genomic Sciences of the HKU. In these workshops, computer science researchers were trained in how to obtain information regarding applications of the life sciences or to provide knowledge of computing for life science researchers. In Hong Kong, the objective of this type of training is to educate both young researchers and instructors. These manpower training and workshop activities were much approved and supported by the Hong Kong SAR government.

The Bioinformatics Special Interest Group (BSIG) brings people working on bioinformatics in Hong Kong together and offers an interdisciplinary environment in which to share information in this field. Group members organise regular seminars and workshops. The Asia Pacific Bioinformatics Network (APBioNET) is the oldest bioinformatics organisation in Asia. It organises the annual International Conference on Bioinformatics (InCoB), which brings together many scientists working in the field of bioinformatics. The sixth InCoB, taking place in Hong Kong on 27–30 August 2007, highlighted the growth in research excellence in the Asia Pacific region [15]. The IEEE International Conference on Bioinformatics and Biomedicine (BIBM) on 18–21 December 2010 and the IAENG International Conference on Bioinformatics (ICB’12) on 14–16 March 2012 were both held in Hong Kong. These conferences provided an international forum for scientists and researchers to exchange scientific ideas in the field.

5 Research in Hong Kong

Research activity in the area of computational biology is increasing day by day in Hong Kong (Fig. 3). Computational biology scientists in Hong Kong are currently inclined towards the areas of database development, sequence analysis and genome analysis. However, other prominent areas of computational biology such as structural bioinformatics, protein networking, gene networking, drug development bioinformatics, systems biology, algorithm development and bioinformatics tools are included in the count of publications shown in Fig. 4a and give a rough indication of the number of publications originating from Hong Kong in those areas. We used the same keywords to find the number of publications in spans of 4 years from 1997 to 2012 and found a significant increase in the number of published papers originating in Hong Kong on different areas of bioinformatics (Fig. 4b).

We have searched research publications in this area in spans of 4 years, i.e. 1997–2000, 2001–2004, 2005–2008 and 2009–2012, using certain keywords. The number of publications provides a rough indication of the expansion in the field. Note, however, that some bioinformatics publications cannot be retrieved using these keywords.

6 Significant Contributions of Computational Biologists

Genome analysis and sequence analysis are the main research priorities in Hong Kong, as is clear from Fig. 4. Significant contributions of Hong Kong scientists working in these areas have been noted. Professor Tsui Kwok Wing Stephen, the director of the Hong Kong Bioinformatics Centre (http://www.hkbic.cuhk.edu.hk/), was involved in the development of the haplotype map of the human genome as a member of the International HapMap Consortium [16]. Research laboratories in the Department of Biochemistry of the CUHK, the Department of Biochemistry and the Applied Genomics Center of the HKUST and the Genome Research Centre of the HKU were involved in the development of the second-generation human haplotype map in which researchers identified more than 3.1 million single-nucleotide polymorphisms (SNPs) [17]. Meanwhile, the Applied Genomics Center and Fok Ying Tung Graduate School of the HKUST, the Department of Biochemistry at the HKUST and the Institute of Digestive Disease at the CUHK participated in cancer genome projects as published in Nature [18]. Human brachydactyly type A1 (BDA1) was the first disorder recorded with autosomal-dominant type abnormalities. A study by Hong Kong researchers revealed that the BDA1 mutation (E95K) impairs the interaction of Indian hedgehog (IHH) receptor PTCH1 with the antagonist HIP1 [19]. Other examples of genome analysis and sequence analysis carried out in Hong Kong are given in Table 2.

7 Tools, Web Servers, Database Research

Hong Kong scientists have also been involved in the development of various bioinformatics tools. Some advanced tools include PriVar, a tool for analysing next-
generation sequencing data, mutations and linkage analysis [20]; COPE, a tool for genome assembly using k-mer frequencies [21]; SOAP3, a rapid graphic processing unit (GPU)-based parallel alignment tool [22]; FetalQuant, a tool for estimating the foetal DNA concentration from maternal plasma DNA which uses maximum likelihood [23]; and GBOOST, a GPU-based tool for detecting gene–gene interactions in genome-wide case control studies [24]. Scientists from the Hong Kong Bioinformatics Centre are also working in this area. The most essential tools developed by these scientists include the ABMapper (a tool for multi-location searching and splice-junction mapping) [25], ViralFusionSeq (a tool for studying soft-clipping, read-pair and targeted de novo assembly to determine and
annotate human viral integration and restructure fusion) [26] and Alns (a tool for searchable and filterable sequence alignment) [27]. Further examples of Web servers or tools developed and maintained in Hong Kong are given in Table 3.

Many bioinformatics databases have been developed and maintained by Hong Kong computational biologists. Some examples are YY1TargetDB (a Web-based YY1 target loci database) [28], BSRD (a comprehensive bacterial sRNAs database) [29] and PcarBase (a searchable database for the brain coral Platygyra carnosus) [30]. Further examples of biological databases developed and maintained in Hong Kong are given in Table 4.

### Table 2  Published work from Hong Kong on genome analysis and sequence analysis by bioinformatics scientists

| Organism type | Gene type/pathogen type | Type of analysis | References |
|---------------|-------------------------|------------------|------------|
| Human         | Locus for type 2 diabetes at 7q32 near PAX4 | Study discovered rs10229583 near PAX4 as a locus for T2D in Chinese and other populations | [49] |
|               | Mendelian disease-causing non-synonymous single nucleotide | Approximate 5% of nsSNVs is pathogenic and carries ~22 pathogenic resulting alleles and this may lead to recessive diseases through consanguineous marriages | [50] |
|               | Loci in or near CDKN1B, TET3, CD80, DRAM1 and ARID5B as associated with systemic lupus erythematosus in Asians | Identified genetic variations in or near CDKN1B, TET3, CD80, DRAM1 and ARID5B which are linked with the disease | [51] |
| MicroRNA-218  | MicroRNA-218 is related to cycle progression and apoptosis in colon cancer. This miR-218 slows down cell cycle progression and endorses apoptosis | | [52] |
| SNP detection | Rapid and precise SNP detection algorithm which help to analyse next-generation sequencing data | | [43] |
| SNP and CNVs at genome-wide scale | Study concluded 79 genes obstruct by CNVs in diseases people and recognised de novo DKK4 duplication | | [53] |
| CD14 gene polymorphism | Study explored the relationship between periodontitis and single polymorphic location in two genes which are DEFB1 and CD14 | | [54] |
| Pathogen      | Salmonella spp. | Study explored occurrence and antimicrobial resistance of Salmonella in meat harvest | [55] |
| Salmonella spp. | Mycobacterium tuberculosis | This experiments conclude about quick detection of Mycobacteria and fast exposure of drug resistance | [56] |
| Mycobacterium tuberculosis | Rat noroviruses | Whole genome sequences | [57] |
| Enterobacter cloacae subsp. cloacae strain ENHKU01 | HIV-1 CRF07_BC variants | Detection of drug resistant mutations | [59] |
| Bacillus macauensis ZFHKF-1 | Structural starting point for RNA binding and homo-oligomer construction | | [61] |
| Influenza B virus | Human coronavirus NL63 | Study deals with disease variety and genetic multiplicity | [62] |

8 Swine Flu, Bird Flu and Respiratory Diseases

Respiratory diseases, as one of the foremost problems in Hong Kong, are a major burden [31]. Respiratory diseases and related viruses are thus important research areas in the territory, and much landmark research has been published in well-known journals such as *Nature* and *Science*. Computational biology researchers are currently investigating the H5N1 virus, strains of which cause H5N1 avian influenza may be transferrable to humans [32]. This virus is highly pathogenic and has caused a pandemic ‘bird flu’ incident in the Hong Kong SAR. Several researchers have reported the emergence of multiple genotypes of the H5N1 virus in terrestrial poultry, leading to an outbreak of avian influenza in chickens in retail markets in Hong Kong and the molecular changes in the virus associated with this event [33]. One study published in *Nature* investigated the long-term evolution and dynamics of transmission of swine influenza A virus (SwIV) using a data set of more than 650 SwIV isolates and more than 800 swine sera from 12 years of systematic monitoring in Hong Kong, including the H1N1/2009 virus that caused a human pandemic [34]. Another study characterised a reassortant progeny of H1N1/2009 with swine viruses [35]. A study published in *Nature* on the origins and evolutionary genomics of swine-
origin H1N1 influenza carried out phylogenetic analysis of the gaps in genetic surveillance and applied evolutionary analysis to estimate the timescale of the origins, finding that a remixture of swine influenza lineages may have occurred years before appearing in humans [36].

9 Collaborative Research

Scientists in Hong Kong are actively collaborating with those in other countries such as the USA, UK, Singapore, Japan, Taiwan and India. Some important research has resulted. One study, a collaboration between researchers in

The table lists examples of biological databases developed and maintained in Hong Kong. The table contains columns for the database name, function of the database, web address, and references.

Table 4 Examples of biological databases developed and maintained in Hong Kong

| Databases name | Function of the database | Web address | References |
|----------------|--------------------------|-------------|------------|
| RedoxDB        | Database for protein oxidative alteration | http://biocomputer.bio.cuhk.edu.hk/RedoxDB | [73] |
| GWASdb         | An spontaneous, multifunctional database for biologists and clinicians to investigate genetic variants and their functional conclusion | http://jjwanglab.org/gwasdb | [74] |
| 16SpathDB      | Database with 16S rRNA gene sequences of all clinically significant bacteria which are listed in the Manual of Clinical Microbiology | http://147.8.74.24/16SpathDB | [75] |
| OpenADAM       | Web-supported data administration system for the large amount of genotype data generated from the Affymetrix GeneChip Mapping Array and Genome-Wide Human SNP Array platforms and it is open source | – | [76] |
| T3DB           | Database that includes bacterial type III secretion process | http://biocomputer.bio.cuhk.edu.hk/T3DB/browse | [77] |
| MMDBD          | An incorporated medicinal resources DNA database | http://www.cuhk.edu.hk/icm/mmdbd.htm | [78] |

the USA, Singapore and Hong Kong, deals with a unique data set arising from surveillance of swine influenza at a Hong Kong abattoir from 1998 to 2010 and may advance understanding of the prevalence of influenza and decrease the occurrence of influenza in Hong Kong [37]. Another study aimed to understand the dissimilarity of two new Dehalococcoides mccartyi strains through collaboration between US and Hong Kong scientists. In this study, Lee et al. [38] performed a comparative genomics analysis via a microarray and concluded that the observed functional incongruence between the activity and core genome phylogenies of D. mccartyi strains is probably caused by a horizontal shift in significant reductive dehalogenase-
encoding genes. Another example of effective collaborative research is the inclusion of different species in the Molecular Ecology Resources Database in which several countries, such as Australia, Austria, the USA, Brazil, Germany, Sweden, Taiwan, China, Canada, India and Hong Kong, participated [39, 40].

The authors of this paper are themselves performing collaborative research. Three of the authors (George, Chakraborty and Zhu) have jointly published research findings on the effects of deleterious non-synonymous SNPs in the binding adaptability of flavopiridol with cyclin-dependent kinase 7 (CDK7), a one-cell-cycle regulatory protein [41]. Another example of collaboration between India and Hong Kong is this review paper concerning computational biology in the Hong Kong SAR; bioinformatics scientists from both countries have critically analysed the activity in and the status of computational biology in the territory.

10 Some Eminent Scientists

Eminent Hong Kong researchers in the field of computational biology include Professor Tsui Kwok Wing Stephen, one of the authors of this review and the director of the Hong Kong Bioinformatics Centre who was a member of the International HapMap Consortium developing the haplotype map of the human genome; two papers on that work were published in *Nature* in 2005 and 2007 [16, 17]. Professor Tsui’s group worked on genomic-sequence variations and the epidemiology of severe acute respiratory syndrome (SARS) [42]. Dr. Jun Wang from the Department of Biochemistry at the HKU is working on computational and transcriptional genomics, structural bioinformatics, SNPs and copy number variation (CNV). He developed an SNP detection algorithm for next-generation sequencing data [43] and also worked on a genome-wide association study on alleles in the FGFR2 gene that is associated with risk of breast cancer [44]. Ascertaining the dynamic nature of gene regulation is a significant challenge in systems biology, an area of research interest in Hong Kong. The corresponding author of this review is working on reconstructing dynamic gene regulatory networks for human cancer, to unravel the dynamic mechanism of cancer development [45]. Dr. Yip Yuk Lap Kevin from the School of Life Sciences at CUHK is researching the use of computational methods to study biological and medical phenomena and networks, and has published work in the journals *Nature* and *Bioinformatics* [46, 47]. Other eminent Hong Kong researchers have been working in the field of computational biology for decades; an attempt to outline the activities of these researchers, to provide a bird’s-eye view of their research activities, is given in Table 5. We have also attempted to determine the proportion of total research in Hong Kong that is related to computational biology, finding that publications in the area of bioinformatics comprise about 9% of total publications (Fig. 5).

The Hong Kong SAR government provides several sources of research funding, such as the General Research Fund (GRF), Collaborative Research Fund (CRF) and Theme-based Research Scheme administered through the Research Grants Council (RGC) (http://cerg1.ugc.edu.hk); the ITF administered through the ITC (http://www.itc.gov.hk); and the Health and Medical Research Fund (HMRF) (formerly known as the Health and Health Services Research Fund—HHSRF—and the Research Fund for the Control of Infectious Diseases—RFCID) administered through the Research Fund Secretariat of the Food and Health Bureau.

The percentage of publications provides a rough indication of the amount of research being carried out. Here, to search for total publications, we used the search term ‘(Hong Kong [affiliation])’. For bioinformatics publications, we used the total number of publications from Fig. 3.

11 Activities of Bioinformatics Companies

Hong Kong relies on medical devices being imported to satisfy the territory’s rising demand for advanced health care. A major focus of the Hong Kong SAR government, therefore, is assisting companies to fulfil the need for medical products. Recent trends in Hong Kong’s population have shown a shift of disease types towards malignant neoplasms, heart disease, pneumonia, cerebrovascular disease, etc. There is a resulting increase in the demand for high-tech products and services that use biotechnology and bioinformatics tools. In response to the outbreak of SARS, many biotechnological companies were established to develop diagnostic kits. One such project involves the development of a biochip by Dr. Yu Cheung-Hoi Albert of the HKUST, which was highlighted in *Science* magazine [48]. This ‘lab-on-a-chip’ rapidly diagnoses emerging pathogens, including influenza viruses; Yu subsequently established the company Hai Kang Life to manufacture these DNA chips. Cluster Technology Limited (ClusterTech) is a computing technology company in Hong Kong which provides solutions for bioinformatics research that combine high-performance computing and cloud computing technology; several other companies have also been established in this area. To assist with world-class infrastructure for such companies, the Hong Kong Science and Technology Parks Corporation (HKSTPC) was established by the Hong Kong SAR government in May 2001 to provide core competency in manufacturing technology, biotechnology, information technology (including computational biology), environmental technology, management systems, etc. The
HKSTPC manages three industrial estates and mentors technology-based companies through its incubation programme which assists with business development for potential new entrepreneurs.

12 Government Role

The Hong Kong SAR government plays a very supportive role in promoting innovation and the development of technology. The government is extremely keen to create an environment which promotes such innovation and development and demonstrates a particular interest in biotechnology, thus fostering favourable conditions for computational biologists. The government supports the innovation and technology support, university–industry collaboration, general support and small entrepreneur research assistance programmes. At the beginning of 2012, for example, 2746 projects were supported from a fund of HK$6.4 billion, with many of the funded projects being related to biotechnology (around 10 %) and information technology (around 20 %), jointly comprising 30 % of the total funding in the territory. The Hong Kong SAR government generally supports fundamental research, applied R&D, technology transfer and technological entrepreneurship, through which it fosters a culture of innovation and a technological environment in the territory.

13 Future Challenges and Concluding Remarks

In the past few years, Hong Kong has seen tremendous growth in bioinformatics research, which has been extended to different fields of biology. However, compared with other developed countries around the world, Hong Kong still has some way to go in this field. Research in the life

Table 5 Some well-known researchers, their affiliations and areas of research interest

| Prominent Scientists/researcher Name | Affiliation | Area of research interest | References |
|-------------------------------------|-------------|--------------------------|------------|
| Professor Stephen Tsui            | School of Biomedical Sciences, The Chinese University of Hong Kong; and Director of the Hong Kong Bioinformatics Centre, Hong Kong SAR | Human cancers and gene regulation, changes in microbiota in human diseases, bioinformatics | [16, 17, 42] |
| Prof. Cheah Kathy                  | Department of Biochemistry, HKU | Working on the directive and function of genes, disease machinery especially skeletal system, human degenerative skeletal disorders and the inner ear | [79, 80] |
| Dr. Yang Wanling                  | Department of Biochemistry, HKU | Working on genomics data assessment and statistical genetics using bioinformatics especially in Asian populations | [81, 82] |
| Prof. Francis Chin                | Department of Biochemistry, HKU | Working on design and analysis of algorithms in the bioinformatics area | [65, 83] |
| Dr. Siu-Ming Yiu                  | Department of Biochemistry HKU | Working on bioinformatics-related algorithms | [84] |
| Dr. Hing-Fung Ting                | Department of Biochemistry HKU | Working on bioinformatics-related algorithms | [85, 86] |
| Dr. Chan Ting Fung                | School of Life Sciences of CUHK | Pursuing research on bioinformatics analysis of microbial genomes and transcriptomes, as well as genetic and genomic analysis of complex human diseases | [87, 88] |
| Dr. Yip Yuk Lap Kevin             | School of Life Sciences of CUHK | Working on using computational methods to study biological and medical phenomena | [46, 47] |
| Prof. Yang Qiang                  | Department of Computer Science and Engineering, HKUST | Algorithms and software tools on computational biology | [89, 90] |
| Prof. Tang Lei-Han                | Department of Physics, HKUST | Working on unravelling the underlying mechanism of conformational transformations of biopolymers as well as modelling the metabolic network of a biological system | [91, 92] |
| Prof. Ng Michael                  | Department of Mathematics, HKBU | Theoretical study on bioinformatics algorithm | [93, 94] |
| Dr. Zhu Hailong                   | Department of Computer Science, HKBU | Hypothetical and computational methods of rebuild gene regulatory networks as well as transcriptional regulatory networks | [45, 95, 96] |
| Prof. Wang Lusheng                | Department of Computer Science in CityU | Working on the algorithms and networks of computational molecular biology | [97, 98] |
Collaborative research activities between bioinformaticians and laboratory biologists at both a national and an international level are also increasing rapidly. Such collaborative research groups analyse biological data ranging from genes to proteins, attempt to solve important biological questions together with laboratory scientists, make various predictions and validate different hypotheses. Despite this positive outlook, however, several challenges exist. More national centres for bioinformation, similar to the National Center for Biotechnology Information (NCBI) in the USA, are needed to provide a central depository and information source for biological data and tools.

During the past few years, development of computational biology has been initiated in Hong Kong through various activities. However, many challenges lie ahead for computational biologists and the Hong Kong SAR government if the territory is to be established as a globally important bioinformatics research centre. The government needs to act to encourage more research projects, teaching modules and conferences and the provision of research grants and support to attract larger numbers to the field of computational biology. Research must be sustained as a key area of investment to improve efficiency and competitiveness in meeting Hong Kong's needs, such as in the medical sector. To identify and support the existing researchers in the field of computational biology, more professional societies and journals indexed in the Science Citation Index should be initiated to drive Hong Kong computational biology towards new heights. This will benefit both national scientists and those in neighbouring Asian countries. Finally, venture capital would be extremely beneficial in the development of bioinformatics companies to assist Hong Kong to become a world-famous centre of computational biology.

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Compliance with Ethical Standards

Conflict of interest The authors declare no conflicts of interest.

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