Preface

Systems biology is a research discipline at the crossroad of statistical, computational, quantitative, and molecular biology methods. It involves joint modeling, combined analysis and interpretation of high-throughput omics (HTO) data collected at many “levels or layers” of the biological systems within and across individuals in the population. The systems biology approach is often aimed at studying associations and interactions between different “layers or levels”, but not necessarily one layer or level in isolation. For instance, it involves study of multidimensional associations or interaction among DNA polymorphisms, gene expression levels, proteins or metabolite abundances. With modern HTO biotechnologies and their decreasing costs, hugely comprehensive multi-omic data at all “levels or layers” of the biological system are now available. This “big data” at lower costs, along with development of genome scale models, network approaches and computational power, have spearheaded the progress of the systems biology era, including applications in human biology and medicine. Systems biology is an established independent discipline in humans and increasingly so in animals, plants and microbial research. However, joint modeling and analyses of multilayer HTO data, in large volumes on a scale that has never been seen before, has enormous challenges from both computational and statistical points of view. Systems biology tackles such joint modeling and analyses of multiple HTO datasets using a combination of statistical, computational, quantitative and molecular biology methods and bioinformatics tools. As I wrote in my review article (Livestock Science 2014, 166:232–248), systems biology is not only about multilayer HTO data collection from populations of individuals and subsequent analyses and interpretations; it is also about a philosophy and a hypothesis-driven predictive modeling approach that feeds into new experimental designs, analyses and interpretations. In fact, systems biology revolves and iterates between these “wet” and “dry” approaches to converge on coherent understanding of the whole biological system behind a disease or phenotype and provide a complete blueprint of functions that leads to a phenotype or a complex disease.

It is equally important to introduce, alongside systems biology, the sub-discipline of systems genetics as a branch of systems biology. It is akin to considering “genetics” as a sub-discipline of “biology”. It is well known that quantitative genetics/genomics links genome-wide genetic variation with variation in disease risks or a performance (phenotype or trait) that we can easily measure or observe in a...
population of individuals. However, systems genetics or systems genomics not only performs such genome-wide association studies (GWAS), but also performs linking genetic variations (e.g. SNPs, CNVs, QTLs etc.) at the DNA sequence level with variation in molecular profiles or traits (e.g. gene expression or metabolomic or proteomic levels etc. in tissues and biological fluids) that we can measure using high-throughput next- and third-generation biotechnologies. The systems genetics approach is still “genetics”, because we are looking at those genetic variants that exert their effects from DNA to phenotypic expression or disease manifestations through a number of intermediate molecular profiles. Hence, systems genetics derives its name, as originally proposed in my earlier article (Mammalian Genome, 2006, 17:548–564), by being able to integrate analyses of all underlying genetic factors acting at different biological levels, namely, QTL, eQTL, mQTL, pQTL and so on. I have provided a complete up-to-date review and illustration of systems genetics or systems genomics and multi-omic data integration and analyses in our review paper published in Genetics Selection Evolution (2016), 48:38. Overall, systems genetics/genomics leads us to provide a holistic view on complex trait heredity at different biological layers or levels.

Whether it is systems biology or systems genetics, the gene ontology annotation is one of the most important and valuable means of assigning functional information using standardized vocabulary. This would include annotation of genetic variants falling into functional groups such as trait QTL, eQTL, mQTL, pQTL. Molecular pathway profiling, signal transduction and gene set enrichment analyses along with various types of annotations form the “icing on cake”. For this purpose, several bioinformatics tools are frequently used. Most chapters in this book and its associated volume cover these aspects.

I would like to point out that systems biology approaches have been proven to be very powerful and shown to produce accurate and replicable discoveries of genes, proteins and metabolites and their networks that are involved in complex diseases or traits. In very practical terms, it delivers biomarkers, drug targets, vaccine targets, target transcripts or metabolites, genetic markers, pathway targets etc. to diagnose and treat diseases better or improve traits or characteristics in animals, plants and humans. In the world of genomic prediction and genomic selection, there have been an increasing number of studies that have shown high accuracy and predictive power when models include functional QTLs such as eQTL, mQTL, pQTL which, in fact, are results from systems genetics methods.

This book and its associated volume cover the above-mentioned principles, theory and application of systems biology and systems genetics in livestock and animal models and provides a comprehensive overview of open source and commercially available software tools, computer programming codes and other reading materials to learn, use and successfully apply systems biology and systems genetics in animals.

Overall, I believe this book is an extremely valuable source for students interested in learning the basics and could form as a textbook in higher educational institutes and universities around the world. Equally, the book chapters are very relevant and useful for scientists interested in learning and applying advanced HTO studies, integrative HTO data analyses (e.g. eQTLs and mQTLs) and computational
systems biology techniques to animal production, health and welfare. One of the chapters focuses on systems genomics models and computational methods applied to animal models for elucidating systems biology of human obesity and diabetes. The two volumes of this book is a result of contributions from highly reputed scientists and practitioners who originate from renowned universities and multinational companies in the UK, Denmark, France, Italy, Australia, USA, Brazil and India. I would like to thank the publisher Springer for inviting me to edit two volumes on this subject, publishing in an excellent form and promoting the book across the globe. I am grateful to all contributing authors and co-authors of this book. I also wish to thank Ms. Gilda Kischinovsky from my research group for proofreading and the staff at Springer involved in production of this book. Last but not least, I wish to thank my wife and children who have given me moral support and strength while I reviewed and edited this book.

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