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Multi-omics data integration considerations and study design for biological systems and disease

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Abstract

With the advancement of next-generation sequencing and mass spectrometry, there is a growing need for the ability to merge biological features in order to study a system as a whole. Features such as the transcriptome, methylome, proteome, histone post-translational modifications and the microbiome all influence the host response to various diseases and cancers. Each of these platforms have technological limitations due to sample preparation steps, amount of material needed for sequencing, and sequencing depth requirements. These features provide a snapshot of one level of regulation in a system. The obvious next step is to integrate this information and learn how genes, proteins, and/or epigenetic factors influence the phenotype of a disease in context of the system.

In recent years, there has been a push for the development of data integration methods. Each method specifically integrates a subset of omics data using approaches such as conceptual integration, statistical integration, model-based integration, networks, and pathway data integration. In this review, we discuss considerations of the study design for each data feature, the limitations in gene and protein abundance and their rate of expression, the current data integration methods, and microbiome influences on gene and protein expression. The considerations discussed in this review should be regarded when developing new algorithms for integrating multiomics data.

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Graphical Abstract

Keywords

Data Integration; RNA-seq; Proteomics; machine learning; microbiome; study design; power

Introduction

The biological system is complex with many regulatory features such as DNA, mRNA, proteins, metabolites, and epigenetic features such as DNA methylation and histone posttranslational modifications (PTMs). Each of these features can be influenced by a disease and cause changes in cell signaling cascades and phenotypes. In addition to the host regulatory mechanisms response to disease, the microbiome can make changes to the expression of the host features such as their genes, proteins, and/or PTMs. In order to gain insight into mechanisms of disease, we need to investigate each of these features and their interplay. For instance, cancers such as melanoma, lung, and thyroid cancers are driven by the BRAF oncogene [1]. However, when patients are treated with therapies that inhibit BRAF, they often develop resistance. Recent multi-omics studies have revealed the heterogeneity and complexity of tumor features such as their genetic mutations, transcriptome, proteins, and signaling pathways. It is now appreciated that tumors can bypass the therapy and give rise to resistance programs [1, 2].

Proper integration of multi-omics approaches has allowed deeper insights into disease etiology, such as unveiling the myriad ways in which the microbiome may play a part in mitigating or enhancing disease risk. This case can be exemplified in regard to incomplete breakdown of bisphenol A (BPA), a mass-produced chemical that is widely used in food packaging, plastics, and resins. BPA has become a growing public health concern as BPA is an endocrine disruptor (as reviewed in Yu 2019 [3]). Thus, research into the fast and complete degradation of BPA, and other compounds via microbial means is of great interest. Yu and colleagues (2019) [3] were able to effectively combine multi-omics data to analyze a microbial community's ability to break down bisphenol A (BPA) products. Though prior research had already discovered the microbes' ability to break down BPA, the interactions

that allowed this reaction were yet unknown. Through a clever multi-omics design, the authors were able to use three major types of integrated analyses to identify differences in encoded and expressed microbial functions that were involved in the BPA-degrading microbial community [3].

Another example, Poore et al. (2020) leveraged multi-omics and machine learning tools, to detect microbial biomarkers from blood and tissues, serving as a great example of microbiome-informed oncology [4]. Here the research team was able to discriminate among healthy and cancer-free individuals as well as between multiple cancer types using plasmaderived, cell-free microbial nucleic acids. Finally, we refer the reader to other reviews about the importance of integrating microbes into multi-omics studies [5–10].

There is a growing appreciation for multi-omics studies in context of therapeutic treatments. However, the methodologies are challenging for a variety of reasons. Each biological regulatory feature has technical hurdles to overcome due to sample preparation, sequencing platforms and depth, limits in instrumentation, and dynamic range [7, 11]. New data integration algorithms are being developed at a rapid pace. In this review, we discuss the background of cellular processes, current data integration methodologies, the considerations for multi-omics study design, and future directions.

Understanding cellular processes in context of 'omics'

Biological systems are complex organisms with many various regulatory features. For instance, the human genome is composed of approximately 3.2 billion nucleotides that give rise to 20,000 to 25,000 protein coding genes, and through alternative splicing events lead to over 1 million proteins (Figure 1). Epigenetic modifications, as well as the microbiome, can influence the expression of both genes and proteins within the biological system under various conditions. In addition to varying numbers of genes and proteins within the biological system, there is also a large dynamic range of high and low abundant molecules within each feature. On top of biological complexity, there are limitations in each of the omic sequencing platforms. These factors should be considered when developing novel data integration methods and are discussed below.

Different organisms have varying numbers of genes and proteins. For instance, there are approximately $4,300, 6,000$, and $25,000$ genes in the E. coli, S. cerevisiae, and H. sapiens genomes, respectively [12]. This leads to approximately 2400 to 7800, 15,000, and 300,000 mRNA molecules per cell for E. coli [13], S. cerevisiae [14], and H. sapiens [15], respectively. Mitochondrial transcripts can account for approximately 20% of polyadenylated RNA. Other high abundant transcripts include those that encode for ribosomal proteins and proteins involved in energy metabolism [16]. It is important to note in sequencing platforms that only a fraction of all transcripts in a sample are actually sequenced and the potentially large number of transcript isoforms generated by alternative splicing events presents another challenge when integrating gene and protein level expression [17]. The transcript isoforms may also change across biological conditions [18]. An overview of the complexity of DNA, DNA methylation, histone post-translational modifications, mRNA, and proteins in humans is depicted in Figure 1.

The estimated number of proteins in a cell is around 2.36 \times 10⁶, in *E.coli* and about 2.3 \times 10⁹ in H. sapiens HeLa cells [19]. Within the vast number of total proteins in a cell, the most abundant proteins can make up 5–10% of protein content and consist of ribosomal proteins, acyl carrier protein (ACP) (functions in fatty acid biosynthesis), chaperones and folding catalysts, proteins of glycolysis (backbone of energy and carbon metabolism), and structural proteins such as actin. Transcription factors are low abundant proteins and range from $1-10³$ copies per cell in bacteria and $10^3 - 10^6$ in mammalian cells. The most abundant proteins usually have many thousands of copies in bacteria and many millions in mammalian cells. The number of genes regulated by a transcription factor depends on its concentration [19]. The protein content depends on the growth conditions and gene induction. Finally, this can become more complicated given the ratio of microbial-to-host cell count, which can depend on host cell type, and other factors [20].

Sequencing technologies for various omics platforms only capture a snapshot of what is happening in a population of cells at one point in time due to limitations in instrument detection, dynamic range, and the lifetime expression of the molecules. For instance, the lifetime expression of mRNA transcripts and proteins are vastly different. The median lifetime of an mRNA in E. coli is 5 min, 20 min in budding yeast, and 600 min for H. sapiens [19]. However, the lifetime of proteins is approximately $1-2$ days. The rate of transcription and translation varies among organisms (E. coli: 10–100 nucleotides (nt) per second (s) and $10-20$ amino acids (aa)/s. H. sapiens: 6–70 nt/s and 2 aa/s; rate of transcription and translation, respectively) (Figure 1). For E. coli a single mRNA transcript can give rise to 10–100 proteins before being degraded. Given this information we can see that there will be an increased chance of detection of proteins with a longer life span, conflating our choice of omics platforms and the resultant interpretations of cellular processes.

It is important to recognize the biological complexity of organisms, dynamic range of molecules, sequencing limitations, as well as the lifetime of expression of those molecules when considering a data integration study design, developing a new algorithm, and when interpreting the results.

Microbiome influences on genes and proteins

In recent years, the importance of the microbiome in host health has been recognized. The idea of the holobiont and the hologenome has had profound implications in how we view the microbiome [21, 22], especially in regard to therapeutics. The idea is that the interactions of the host's own genome and its "second genome" [23], collectively called the hologenome, work together to provide an "insurance policy" against a variety of perturbations [24, 25] that affect host health. This close relationship of microbe-host interactions can be more explicitly termed the "microbiota-nutrient metabolism-host epigenetic-axis" [26].

Microbiota and their metabolites can affect the host epigenetic landscape, by directly modifying histones, altering DNA methylation profiles, and influencing the nature of noncoding RNAs (Figure 1). For example, histones can be modified by microbiota by

altering the activity of histone modification enzymes, and the levels of the enzymes substrates [27–29].

Microbiota can also affect the therapeutic nature of drugs. Many prodrugs, i.e. a drug that must metabolically converted in order to become pharmacologically useful, may remain inactive (i.e. the microbiota that mediate the conversion of the prodrug to its active form are not present), or the drug / prodrug, may not become bioavailable to the host, as a result of degradation by the hosts microbiota [30]. Moreover, patients taking NSAIDs (non-steroidal anti-inflammatory drugs), may promote the preponderance of antibiotic resistant bacteria as 24% of tested over-the-counter NSAIDs inhibited the growth of at least one microbe in vitro [31]. These metabolomic effects, raises concerns about potential side-effects of therapeutic drugs, or other diet and treatment regimens, intended to be used on humans and agricultural systems. For example, antibiotics can eliminate histone deacetylase (HDAC) inhibitorproducing microbes. These microbes, when present, can augment regulatory T (Treg) cells, which aids in anti-inflammatory processes [32].

This means that the diversity of microbial metabolic pathways, and their impact on drug pharmacokinetics and pharmacodynamics [33, 34], may partly explain the variation to drug responses between individuals and populations. Therefore, therapeutic treatments that involve the microbiome, may have to be regionally tailored [30, 35, 36].

Histones can undergo both variant replacement and post-translational modification (PTM), together these form the "histone code". These local arrangements can affect chromatin structure in such a way that leads to the activation or repression of transcriptional activity [37, 38]. Thus microbes, through diet, have the ability to modify methylation and PTM profiles of the host, and can also affect the generation of short-chain fatty acids (SFCAs) through the fermentation of dietary carbohydrates. SFCAs, such as butyrate and acetate, can inhibit deacetylase levels. Meaning that chromatin structure becomes increasingly relaxed due to acetylation driving increased transcriptional activity [26]. In fact, it has been shown that microbes can affect host tissue acetylated and methylated chromatin states in a sitespecific and combinatorial fashion and even impact host developmental and metabolic phenotypes [37–39].

Modelling the development of the microbiome and its commensurate ontogenetic changes of the host, are increasingly being considered when trying to interrogate host health and therapeutics [40]. Many microbial ecological principles such as community assembly are being brought to bear to investigate these processes [41, 42]. These changes can be exemplified through host immune maturation, considering that the host immune system must not only be able to recognize "self" antigens, but also those of symbiotic microbes. How microbes influence the expression of the major histocompatibility complex (MHC), or how host heterozygosity in turn affects the diversity of the microbiota through MHC, is largely unknown and is an active area of study [26, 43]. Finally, the role of microbes as they relate to cancer and immune treatments are increasingly becoming targets for the development of therapeutic strategies [44, 45].

Proteomics, in combination with other omics strategies have been used to interrogate disease processes. However, if we do not take into account the effects of microbiota (i.e. the entirety of the holobiont), then we may miss meaningful insights to develop potentially therapeutic treatments. Particularly those related to metabolic disorders (e.g. obesity), or the systemic effect of metabolites (e.g. bile acids) on organ systems [46]. There is far more variation of our "second genome" that can be leveraged for human benefit compared to our own [47].

Advances in Microbial Ecology

With heavy emphasis on understanding the effect of the microbiome it has become common practice for biomedical researchers to include methods to investigate the diversity of bacteria and archaea in their samples. The history of microbial ecology centers around the sequencing and alignment of appropriate phylogenetic marker genes. The 16S rRNA gene, first purposed as a marker by Woese and Fox (1977) [48], is by far the most commonly used marker gene with massive databases of full length gene isolate from environmental and culture-derived sources (e.g SILVA, RDP, Greengenes [49–51]. (Table 1). New microbial taxonomy databases, such as the Genome Taxonomy Database (GTDB), not only curate a 16S rRNA gene reference database, but are also leveraging phylogenomic information [52, 53] to provide a consistent framework for determining the phylogenetic context partial or complete genomes derived from metagenomes [54].

Apart from selecting a marker gene and appropriate database, researchers also have a choice between sequencing methods and platforms. Due to limitations of short-read platforms such as Illumina and Ion Torrent, researchers must select between variable regions of the ~1500 bp 16S rRNA gene. Each variable region provides a different level of sensitivity and specificity depending on microbial community composition. This is why preliminary amplicon surveys often compare a collection of primer sets and variable regions. The combination of the primer set and the amplicon region that best differentiates among the common taxa in the study, is then chosen. Alternative long-read platforms have recently been adapted to deliver high-throughput full-length 16S rRNA for researchers that need taxonomic resolution beyond the genus to family level typically provided by short-read technologies [55].

Current metagenomic analysis techniques have allowed researchers to obtain partial and complete draft genomes from environmental/host-derived samples given sufficient sequence coverage. This coverage factor is highly dependent on the species evenness and richness. Researchers using these techniques can investigate potential functional differences of a collection of metagenome assembled (draft) genomes. However, often they have to use concentrated universal proteins to place these genomes in a phylogenetic context because of the difficulty of assembling and correctly binning highly conserved genes like the ribosomal subunit genes. Combined universal marker genes are used to construct the phylogeny from genomes assembled from environmental and host-derived sequences along with a minority of familiar microbial genomes from culture collections. The sudden rush of sequencing microbial genomes has necessitated the construction of easy-to-use wrappers and pipelines to aid biologists in learning how to approach the analysis of their metagenome data, either in whole or in part. Some great examples of such tools are, QIIME 2 [56], metaWRAP [57],

Sunbeam [58], SqueezeMeta [59], metAMOS [60], mg-RAST [61], IMG/M [62], Anvi'o [63], MicrobiomeAnalyst [64], and the variety of tools within the biobakery [65] collection (e.g. MetaPhlan2 [66], PhyloPhlan [67], HUMAnN [68], LEfSe [69]), among others (see [70] for a review these and many other meta'omics tools). Biomedical researchers wading into the depths of microbial ecology looking to integrate disease metrics, host proteomics, and microbial diversity should be aware of the various databases, curatorial rigor, and the limitations of the sequencing platform they choose.

Sequencing technologies

Depending on the biological question, there are many types of omics technologies targeting DNA, total RNA, mRNA, miRNA, DNA methylation, proteins, protein modifications, histone post-translational modifications, metagenomics, metaproteomics, etc. Sequencing platforms have improved over the years and now allow for the sequencing of large complex human samples within a few days from small amounts of material (Table 2). Several workflows have been developed to sequence the whole genome, the whole exome (proteincoding portion of DNA), and transcriptome (mRNA), and arrays for specific cancer or immune-related genes. In addition, we can profile modifications, such as DNA methylation using either whole genome bisulfite sequencing or Illumina's MethylationEPIC BeadChip arrays. The detection of such modifications can also be determined through the direct sequencing of long read DNA and RNA via the Oxford Nanopore Technologies (ONT) MinION platform [71–74], and PacBio instrumentation.

Error rates and read lengths vary between DNA sequencing technologies. Illumina short read sequencing (i.e. Hiseq, Miniseq, etc.) typically have very low error rates, at about .25% per base, but are sensitive to low diversity libraries, as is the case with applications such as 16S metagenomics and targeted gene approaches. Long read technologies have higher error rates, ranging from 13–15% for PacBio and 5–20% for Oxford Nanopore instruments [75, 76]. Read length for Illumina platforms have a maximum length of 600 bases but long read technologies commonly achieve 10–30kb for a single read [77]. Optimal read length is also dependent on the application. Where most sequencing experiments can collect suitable information with 150 – 300 base pair read lengths, there are exceptions. Illumina's 16S Metagenomics protocol requires 2×300 base pairs. For whole genome sequencing (WGS), the longest read possible is optimal but with long read technologies, the error rate increases with the length. Many researchers have combined long read and short read sequencing to "fill gaps" with WGS. Due to the fairly recent advent of long read sequencing technology, information on optimal long read lengths for applications other than WGS is sorely lacking but Illumina short read sequencing is rich in optimal read length recommendations [78].

Long read sequencing technologies such as ONT and PacBio have already ushered in significant improvements in both the amplicon and metagenomic sequencing space. From high resolution analysis of the full length 16S gene [55], the entire rRNA operon, [79], to improving the ability to close entire microbial genomes [80]. For an in-depth overview on these long-read sequencing technologies, see Amarasinghe et al. [77].

Mass spectrometers have also improved by increasing sequencing depth capabilities over the past 5–10 years. The technology has advanced from sequencing roughly 3,000 proteins in a cell line experiment using older LTQ mass spectrometers to routinely sequencing 8,00010,000 proteins using newer Orbitrap Lumos and Orbitrap Eclipse mass spectrometers. Most proteomics experiments are performed using data dependent acquisition (DDA) mode. In this method, the top 20 most abundant peptides in the MS1 scan that are eluted from a liquid chromatography (LC) column are selected for fragmentation in the orbitrap in order to generate the peptide sequence MS2 scan. The complexity of the sample mixture highly influences the sequencing depth and how many proteins will be identified. Understanding the protein abundance and make-up of the samples is critical. If transcription factors are the target molecules, then some method of removing highly abundant proteins prior to mass spectrometry may be necessary. This is especially critical for serum and plasma samples that have high abundant molecules, such as albumin and hemoglobin. Otherwise, the mass spectrometer will sequence thousands of molecules of albumin and miss the most interesting low abundant proteins [81].

The latest mass spectrometry technology utilizes data independent acquisition (DIA) to sequence all of the peptides from the MS1 scan as they elute from the LC column as opposed to DDA methods that only sequence the top most abundant peaks. DIA methods are beneficial over DDA for complex mixtures, such as in the serum example above. This method helps to overcome complex mixtures that are highly influenced by high abundant proteins [82–84].

In addition to shotgun sequencing for the host genes and/or proteins, we can also utilize shotgun sequencing for the microbiome. Shotgun metagenomics/metaproteomics may only sample the dominant microbiota when the sequencing depth is very shallow. A major challenge of shotgun sequencing the microbiome is the difficulty in assembling genome fragments due to under sampling, it is also just as difficult to piece together peptides for robust protein and taxa identification.

Despite these potential issues, it is possible to sample the microbial proteome in depth from a variety of human body sites and diseases, such as saliva, gut / feces, cervicovaginal, or chronic kidney disease [40, 85–87]. However, the study / sampling design and analytical approaches one must consider can differ greatly between each study. Several sampling preparation approaches have been shown to enrich microbial biomass ranging from differential centrifugation through double-filtering differential separation. These approaches are often followed by a variety of optimized microbial lysis protocols, typically involving mechanical disruption (e.g. bead beating, sonications), complemented with enzymes (e.g. trypsin) and detergents. Upon successful lysis, it is just as important that remaining enzymes, detergents and salts be removed. For more details see the review by Issa Isaac et al. and Lin et al. [7, 40] and the references therein.

Another complication for metaproteomics experiments is due to the fact that proteins within the same organism have shared peptide sequences. In order to have confidence in the protein identification, a unique peptide match for the protein should be identified with high confidence. This is made even more complicated when mapping peptide sequences to

hundreds of different species that have conserved protein sequences. Mass spectrometry does not sequence proteins, but rather measures the mass-to-charge of peptides and relies on mass spectra matches to a database of protein sequences for protein identification. However,

Curated databases are critical to properly analyze nucleotide and protein sequencing data generated from these various sequencing platforms. The ability to align reads to a reference genome is only as good as the sequence and annotation information present in the reference genome. There are several resources that continually curate and update nucleotide sequence information and annotation including University of California Santa Cruz (UCSC) Genomics Institute genomes, National Center for Biotechnology Information (NCBI) GenBank and RefSeq, Encyclopedia of DNA Elements (ENCODE), and Ensembl to name a few. The Universal Protein Resource (UniProt) contains both Swiss-Prot (manually annotated and reviewed) and TrEMBL (automatically annotated and not reviewed) databases for protein sequence information (Table 1).

there is hope to make sense of these (Table 1 and Table 2) [7, 40].

Data Integration and current methodologies

Several data integration methodologies have been developed to integrate certain types of omics data. In addition, large data repositories have been created to house data from sequencing experiments for various diseases. These resources provide valuable building blocks and large amounts of biological samples that can be utilized to push data integration methods forward. Currently, data integration tools implement a variety of methods but generally fall under two categories: multi-staged analysis and meta-dimensional analysis [88]. Multi-staged integration models are constructed using only two numerical or categorical features of the data. For example, gene counts from an RNA-seq experiment are combined with protein information from a mass spectrophotometry run. Meta-dimensional analysis attempts to incorporate all the types of data of interest by concatenation or transformation into a simultaneous matrix or "metadata" set that can be analyzed simultaneously. The latter method has more statistical power but can be challenging when attempting to combine data from different types of datasets. Yet, how does a researcher decide which tool or method is most appropriate? As stated above, the biological question is the driving force in the type of analysis method chosen and factors such as sampling, the type of platform, and quality of the data are important. How were the samples collected and prepared? Can the data be effectively analyzed if sequencing depth or quality is low? Are the data types compatible? How much signal is lost after normalization and filtering? These are all questions that should be considered before choosing the appropriate tools.

Unfortunately, data integration and analysis are very complicated and there currently do not exist many user-friendly tools for researchers with limited bioinformatics backgrounds. Many tools utilize the statistical language R, which requires programming expertise in addition to strong biostatistical knowledge. For example, the R package integrOmics, which combines proteomics, transcriptomics, and pathway analysis on two data sets uses correlation analysis and partial least squares regression [89]. The R package mixOmics uses multivariate analysis for data exploration, dimension reduction and visualization [90]. Micrographite integrates miRNA and gene expression with pathway analysis [91] and

iClusterplus [92] and LRACluster [93] use clustering to integrate methylation and gene expression data (Table 3).

For both multi-state and meta-dimensional methods, many different algorithms are used, but the most common ones are clustering, network analysis, data reduction (PCA), and Bayesian analysis [94]. Ray et al (2014) used Bayesian analysis to analyze gene expression and methylation data in ovarian cancer using data collected from the Cancer Genome Atlas Project and detected a gene, SPON1, which appears to be regulated by methylation of its CpG site [95]. Correlation based analysis can be useful when prior knowledge of biochemical interactions is lacking [96]. Regardless of the methodology, appropriate normalization and data filtering is very important as data is being incorporated from multiple sources.

There also exist some web-based tools such as Paintomics [97] that attempt to make the data analysis easier but can still be difficult for the inexperienced user and the researcher must have a good working knowledge of their data [98]. Further, there are databases that are commonly used in integrated omics analysis, such as the Cancer Cell Line Encyclopedia (CCLE), The Cancer Genome Atlas Program (TCGA), Tumor Alterations Relevant for Genomics-driven Therapy (TARGET), and Omics Discovery Index (OmicsDI) [94] (Table 1). CCLE and TCGA have characterized thousands of cancer data sets and can be used for data mining and visualization. TARGET utilizes clinical information and has resources for analytical tools on their websites. OmicsDI provides a platform for searching public and protected data for a large variety of organisms.

Considerations for study design and power evaluation

As for any high-quality study, conducting a multi-omics study should always begin with identifying the scope and restrictions of a study. Careful planning and execution will improve a study's robustness and reproducibility and are especially crucial in multi-omics studies, as they involve a large number of comparisons, tailored statistical analysis, substantial financial and timely investments [10, 98]. Involving a statistician from the very beginning of a study is critically important to assist the researcher to identify the research question, define clear a priori hypotheses, proper experimental design, study analysis and interpretation, drawing conclusions and much more [99, 100].

Once research hypotheses are clearly defined, a suitable study design is selected that addresses the research hypotheses best. Therefore, several questions need to be evaluated, such as: Are one or more intervention groups compared to a control (or themselves), or is an effect evaluated in the same samples before and after intervention? Is an intervention effect over one period of time or will samples be measured at several different time points? Will biological samples be pooled or analyzed individually and what is the scientific justification for it [101]? Which types of omic platforms will provide the most value [101] and how are the multi-omics data going to be integrated? Are samples from the same biological source available for all multi-omics platforms of interest? Ideally, samples for all omic platforms would be collected from the same source. However, this is not always possible due to sample-specific limitations or accessibility and amount of the material [101]. For instance,

generating multi-omics data from formalin-fixed paraffin-embedded (FFPE) tissue might not be possible for certain omic platforms [101]. While there are many questions to be considered during the selection of the experimental design, the deciding factor for the choice of a study design is usually its feasibility and financial limitations [10, 102].

Following the selection of a study design, available resources need to be allocated between the individual omic platforms [101]. This allocation should be guided by the cost and contribution of each individual omic platforms to the multi-omics study as well as the statistical power of each individual omic platforms. Omic platforms with a substantial signal-to-background noise ratio will require less samples and allow for an allocation of more resources to omic platforms with a small(er) signal-to-background noise ratio, as these platforms require more samples to achieve (similar) adequate statistical power. In addition, some omic platforms will also require some internal distribution of resources. For example, when designing an RNA-Seq study the trade-off between the number of samples and sequencing depth will need to be balanced [10].

The sample and data collection should be guided by the data analysis to reduce confounding and technical artifacts, such as batch effects [10]. These effects can be introduced during steps of the sample collection, preparation and storing (e.g. multiplexing) [10, 101]. While some ad hoc methods attempt to reduce such biases introduced by technical artifacts, they are inferior to a randomized design [98]. However, some technical artifacts cannot be avoided, and in these cases it is important to identify and understand such limitations early in experimental design to mitigate and recognize their impact on the results and conclusions [10, 101].

Due to the complexity and large volume of data associated with multi-omics studies it is crucially important to tailor the statistical analysis to a specific research project [10]. A variety of methods for integrating multi-omics data have been proposed and categorized as either supervised, semi-supervised, or unsupervised [103]; as well as, conceptual, statistical, correlation, network, and model-based integration [102]. The integration and statistical analysis of a multi-omics study depend on the selection of omic platforms and their associated types of data (e.g. count values, percentages). Nevertheless, each analysis and method have its underlying assumptions that need to be verified [98].

As in any well-designed study, an initial power calculation is increasingly crucial to evaluate and estimate a sufficient number of samples and avoid a potential waste of resources, especially in such large-scale studies [10]. Power is defined as the probability of correctly rejecting the null hypothesis, which is the likelihood of detecting a true signal or effect. A mathematical power calculation is usually impossible due to the complex nature of the study design and data; however, estimating statistical power using simulation is a valuable alternative. The evaluation of statistical power involves four major steps. First, data needs to be simulated and a pilot study, prior knowledge, literature or experts can be beneficial for the generation of realistic data. Next, a true signal needs to be introduced and should be guided by the expected effect size. The data can then be analyzed, and the statistical power can be estimated empirically. In the context of a case-control study, the statistical power is the percentage of correctly identified features out of all features with an introduced effect size.

The statistical power of a study depends on several factors (Figure 2) of which some can be controlled, while others are fixed due to the study and its design. First, the choice of statistical method used for the analysis. While some tests are more powerful than others, it is important that their assumptions are verified and met. Another factor that influences the statistical power of study is the number of variables measured by the individual omic platform, which is usually dictated by the omic platform [101]. For example, genomics typically measures millions of variants [104, 105], transcriptomics quantifies tens of thousands of molecules [106], and proteomics [107] and metabolomics [108, 109] profile thousands of molecules. Further, statistical power is affected by the magnitude and prevalence of the effect of the phenotype or exposure (effect size). How distinct is the effect? How substantial is the signal difference between groups? And how many measured variables are affected? Information about the effect size might be available from previous literature or expert knowledge but is often unknown [101]. In such cases, a pilot study can assist with estimating the effect size, but these estimates need to be handled with caution due to instability [101, 110]. Yet another power influencing factor is the homogeneity of the measured values, describing the natural variance of the sample, the precision of the measurement instrument and detection limits. With an increasing variance the statistical power will be reduced. The variance of the samples can be the result of many aspects, such as the sample population selection, choice of tissue type or confounding factors [111]. In addition to sample variance inflation, confounding factors can also introduce biases in the data, and therefore it is important to collect sample meta-data to mitigate some effects of confounding [101]. Because most of the factors affecting the statistical power of a study are fixed or dictated by the study design, the factor that is most commonly used to adjust the statistical power of a study is the sample size.

Applying power analyses for microbiome data is still a burgeoning field of inquiry and is replete with difficulties [112–115]. The types of power analyses, like those outlined above, differ based on the questions being asked of microbiome data. Typically, power analyses of microbiome data center on measures of alpha and beta diversity, and differences in compositional abundances of taxa [114, 116, 117]. Which of these measures to use will depend on the question at hand. How to integrate these into a multi-omics study is still underdeveloped [115].

Power and sample size evaluation is a valuable technique during the experimental design of a study to ensure adequate power and sample size. While under- and overpower studies unnecessarily deplete resources, the risk of failure of a study is especially prevalent in underpowered studies. Underpowered studies and studies with improper experimental design are more likely to miss true signals, produce bias results, false positive (type I error) and false negative (type II error) results, which will lead to misinterpretations [10, 101, 111]. Such incorrect inferences will impact the reproducibility, scientific progress and the cost of science [98, 118].

Conclusion and Future Directions

It is important to consider the context of the disease or research question that is under investigation and what types of data will provide valuable insight when integrated together.

Depending on the biological question, type of material (fresh tissue, FFPE tissue, serum/ plasma, and cell lines), amount of DNA/RNA/protein, number of biological replicates, and the number of confounding effects in a study, these factors will determine the best sample preparation and sequencing methods needed for data acquisition. Sample preparation methods including the day each sample is prepared, the type of DNA, RNA, and/or protein that is extracted, library generation for genomics, protein digestion and peptide labeling methods for mass spectrometry, and the sequencing platform/instrumentation are all key factors in the study design and the interpretation of the final results. If one sample is prepared on a different day than the other biological replicates, this will introduce variance and/or bias and reduce the statistical power of the analysis. If proteomics samples are multiplexed using multiple TMT-10plex batches, this will introduce a batch effect across sequencing runs. These factors should be discussed prior to sample preparation.

It is also critical to know what population of regulatory features were captured for sequencing and can be integrated. For example, membrane bound proteins cannot be integrated with gene expression data if membrane proteins were not solubilized during sample preparation prior to performing mass spectrometry. A caveat with mass spectrometry data is the fact that a missing value does not necessarily mean a protein is not expressed, but only that the protein is below the detection limits of the mass spectrometer. The biological questions should be a driving force in the methodology used for multi-omics data integration.

Though multi-omics datasets can provide an individual with a greater depth of understanding in certain scenarios, this is not without cost. Omics studies often rely on large numbers of comparisons, the correct data type, appropriate statistical analyses, and a considerable investment of time, skilled personnel, and money. When constructing an experiment one must be weary of what types of omics data can and should be integrated to achieve the greatest understanding of the system being studied [98]. High throughput omics platforms are not always necessary to answer the research question. Traditional techniques, such as enzyme-linked immunosorbent assay (ELISA) assays, immunohistochemistry (IHC), and quantitative polymerase chain reaction (qPCR), may be all that is necessary to validate a particular biological mechanism. In fact, these techniques are often required to validate the findings from a larger omics study in order to verify the significant molecule identified from omics data is a true positive result.

For the most part, current tools utilize clustering, networking, data reduction and Bayesian analysis. Because of ever increasing acquisition of data, resulting in large datasets and increasing numbers of them, machine learning will become more and more necessary for effective analysis and data mining. There is a need for accessible and well documented methods, tools and algorithms [96]. As with all scientific endeavors, the easy questions will be answered first and "low hanging fruit" will be become less prevalent. Thus, there is a need for more effective algorithms and computing resources [88]. Because of the variety of platforms used to generate multi-omics data, standardization of data formats would make integration easier [94].

Future multi-omics data integration algorithms should take advantage of the "big data" resources (Table 1) and the advent of machine learning and artificial intelligence algorithms [10, 119, 120]. Machine Learning has played an increasingly important role in allowing scientists to integrate multi-omics datasets. By utilizing a machines ability to compare and identify patterns in large quantities of biological data, we allow for far more accurate and efficient methods of elucidating complex cellular mechanisms and in some cases providing predictions to clinical outcomes. This is achieved through the computer's unique ability to observe multiple layers of omics data simultaneously providing a more holistic view of the systems at play, rather than observing each omic system individually and drawing simple conclusions based on visible correlations [4, 120].

New data integration methods should include variables related to each omic platform's weaknesses and limitations. Each method is limited by its statistical power, sample size, technical variables, batch effects, sequencing depths, sample preparation, and a multitude of other factors. These factors are important to keep in mind when designing, conducting and analyzing a study and interpreting the results. Therefore, it is highly recommended to involve a biostatistician/bioinformatician from the very beginning of any study. Their expert knowledge can be valuable at any stage of a study to prevent errors, wasting resources and optimize the study. The need for trainings program in this rapidly evolving field has been recognized by many institutes, such as Jackson Laboratory, [Bioinformatics.org,](http://Bioinformatics.org) UC Davis and Johns Hopkins, and many bioinformatics training programs are available online for free or with costs. Lastly, researchers should always remember to validate significant findings using other traditional wet lab techniques to unmask false positive results.

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Figure 1. Overview of chromatin structure and gene/protein regulation.

DNA access is regulated by DNA methylation and histone post-translational modifications (PTMs). There are approximately 3.2 billion nucleotides in the human genome transcribed to approximately 20,000–25,000 protein coding genes, which are translated to over 1 million proteins due to alternative splicing events. Each layer of regulation can also be modified by microbes that are present in the environment and host organism. Each level of biological regulation can be sequenced by using various nucleotide and protein/peptide sequencing technologies.

Figure 2. Factors that influence the statistical power in multi-omics studies.

The statistical power of a multi-omics study can be effected by several factors including and must be considered at the beginning of the study. Such factors include, but are not limited to (the effect of the following factors on power are under the assumption that the remaining factors remain constant): (1) The type of the study. While randomized controlled studies are generally more powerful than observational studies due to controlling unwanted effects, limitations can prohibit this application of a randomized controlled study. (2) The sample allocation. In general, a balanced study, where samples are equally distributed among group, is more powerful unbalanced study. (3) Sample size. As the number of samples in a study increases the statistical power improves. (4) Effect size. The greater the true differences between groups, the greater the statistical power of a study. (5) Hypothesis test. While parametric tests are in general more powerful than nonparametric test, parametric tests are not applicable if there assumptions are not met. (6) Significance level α . The significant level represents the probability of type I errors, the probability of rejecting the null hypothesis given that the null hypothesis is true. As the numerical value of α increases, the probability of type I errors increases as well as the statistical power (probability of rejecting the null hypothesis given that the null hypothesis is true). (7) Number of tests. Testing multiple hypotheses requires a correction and reduces the statistical power. (8) Background noise and sample variation increase the variance and complicate the detection of a true signal and therefore decrease the statistical power. (9) Confounders can increase variance and/or introduce a bias, which decreases the statistical power.

Table 1.

Available resources for big data sets.

The table list resources available to download data sets from various omics platforms as well as sequence and annotation information.

Table 2.

Coverage and Read Recommendations by Application.

Each genomics platform has a recommended sequencing depth depending on the biological question and focus [136].

Table 3.

Data integration tools for multi-omics.

Algorithms and bioinformatics tools for data integration across multiple omics platforms.

