Proceedings of the fourth international molecular pathological epidemiology (MPE) meeting

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Abstract
An important premise of epidemiology is that individuals with the same disease share similar underlying etiologies and clinical outcomes. In the past few decades, our knowledge of disease pathogenesis has improved, and disease classification systems have evolved to the point where no simple disease processes are considered homogenous. As a result, pathology and epidemiology have been integrated into the single, unified field of molecular pathological epidemiology (MPE). Advancing integrative molecular and population-level health sciences and addressing the unique research challenges specific to the field of MPE necessitates assembling experts in diverse fields, including epidemiology, pathology, biostatistics, computational biology, bioinformatics, genomics, immunology, and nutritional and environmental sciences. Integrating these seemingly divergent fields can lead to a greater understanding of pathogenic processes. The International MPE Meeting Series fosters discussion that addresses the specific research questions and challenges in this emerging field. The purpose of the meeting series is to: discuss novel methods to integrate pathology and epidemiology; discuss studies that provide pathogenic insights into population impact; and educate next-generation scientists. Herein, we share the proceedings of the Fourth International MPE Meeting, held in Boston, MA, USA, on 30 May–1 June, 2018. Major themes of this meeting included ‘integrated genetic and molecular pathologic epidemiology’, ‘immunology-MPE’, and ‘novel disease phenotyping’. The key priority areas for future research identified by meeting attendees included integration of tumor immunology and cancer disparities into epidemiologic studies, further collaboration between computational and population-level scientists to gain new insight on exposure-disease associations, and future pooling projects of studies with comparable data.

Use of Standardized Official Symbols For unambiguous communication, we use HUGO (Human Genome Organisation) Gene Nomenclature Committee-approved official symbols (or root symbols) for genes, gene products, and gene families, including APC, AR, AXIN2, BRAF, CD4, CD8, CTLA4, CTNNB1, EGFR, ERBB2, ERG, ESR1, FOXA1, FOXP3, HLA, IDH1, IDH2, IGF1, IGF1R, IGFBP, Kras, Muc1, MKI67, PDCD1, PGR, PTEN, RB1, TERT, TMGRS5, and Wnt, all of which are described at www.genenames.org. Each colloquial name is used in parenthesis following its official symbol counterpart. This format enables readers to familiarize themselves with the official symbols for genes and gene products together with common colloquial names.

Extended author information available on the last page of the article
**Keywords** Molecular pathological epidemiology · Meeting report · Meeting proceedings · Meeting summary · Patho-epidemiology

**Abbreviations**
AI  Artificial intelligence  
AICR  American Institute for Cancer Research  
BCAC  Breast cancer association consortium  
BCR  B cell receptor  
CDR  Complementarity determining region  
CIMP  CpG island methylator phenotype  
cfDNA  Cell free DNA  
ctDNA  Circulating tumor DNA  
ddPCR  Droplet digital PCR  
eQTL  Expression quantitative trait loci  
FFPE  Formalin-fixed paraffin-embedded  
GBM  Glioblastoma multiforme  
GWAS  Genome-wide association study  
HPFS  Health professionals follow-up study  
IHC  Immunohistochemistry  
MPE  Molecular pathological epidemiology  
MSI  Microsatellite instability  
NHS  Nurses’ Health Study  
PCR  Polymerase chain reaction  
PRS  Polygenic risk score  
RTK  Receptor tyrosine kinase  
SNP  Single nucleotide polymorphism  
TCGA  The Cancer Genome Atlas  
TCR  T cell receptor  
TMA  Tissue microarray  
WCHS  Women’s Circle of Health Study  
WCRF  World Cancer Research Fund  
WES  Whole exome sequencing

**Introduction**

A central objective of epidemiology is to investigate why some diseases (or related health events) occur in certain groups of people but not in others, with the view to applying that knowledge to prevent or control those outcomes in the future. Traditionally in this effort, diseases or health events are treated as binary outcomes; for example, a group of people either have or do not have cancer at a given organ site and, if etiology is of interest, rates of that outcome can be compared across levels of a potential exposure. While this approach has identified major causes of morbidity and mortality in human populations that have subsequently supported important public health policy changes, the over-simplified ‘yes vs no’ nosology misses the essential concept of pathogenic heterogeneity. Most human diseases, and essentially all cancers, are biologically different from one patient to the next. Indeed, many human diseases are complex processes that occur for reasons that are, in a precise sense, unique to that individual, the result of that person’s specific host characteristics (e.g., genome) and a multitude of distinct external factors (e.g., diet, lifestyle, environmental, microbiome). Molecular pathological epidemiology (MPE) is an integrative scientific discipline that examines the interplay of these unique disease, host, and external factors.

One of the main aims of MPE is to investigate potential etiologic/survival factors across strata of molecular characteristics for the disease-of-interest. The underlying premise with an etiologic study in MPE is that diseases that share certain molecular alterations are more likely to share common causes; similarly, for prognosis studies, the general hypothesis is that some external or endogenous factors may influence disease outcomes according to molecular attributes because those factors likely interact with the diseased cells in the local tissue microenvironment. Historically, these molecular classifications were often drawn from the clinic, such as microsatellite instability (MSI) for colorectal cancer and statuses of ESR1 (estrogen receptor 1, ER), PGR (progesterone receptor, PR), and ERBB2 (HER-2) for breast cancer [1–6]. With the recent growth of high-throughput biologic data in large epidemiology studies, disease phenotyping has become more sophisticated and includes tumor sequencing, gene expression, proteomics, and epigenomics.

The International MPE Meeting Series began in April 2013 as a small, local meeting of 10 investigators at the Harvard School of Public Health. Subsequent meetings became larger, with over 150 scientists from more than 16 countries attending each of the Second (December 2014) [7] and Third (May 2016) [8] International MPE meetings. Because MPE is inherently transdisciplinary and it is a relatively new scientific discipline, these meetings gave attendees a rare opportunity to share ideas, methods, successes, and challenges; further, they were an opportunity to help train the next generation of MPE scientists. Herein, we share the proceedings of the Fourth International Molecular Pathological Epidemiology (MPE) Meeting, held in May/June 2018 at Dana-Farber Cancer Institute in Boston, MA, USA. A list of the speaker names, lecture titles, and key references appears in Table 1.

30 May 2018

For the first time in the history of the International MPE Meeting Series, a pre-meeting interactive workshop was held for current and future leaders in MPE. Dr. Reiko Nishihara chaired the session which included panelists from a
Table 1  Summary of podium presentations at the 4th international molecular pathological epidemiology (MPE) meeting in Boston, MA, on 31 May and 1 June, 2018

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wide range of career stages and institutions (Drs Christine Ambrosone, Peter Campbell, Montserrat Garcia-Closas, Marios Giannakis, John Quackenbush, and Molin Wang). The workshop consisted of questions from the session chair and audience members with responses from panelists concerning grant writing for transdisciplinary science, transdisciplinary team building, and training opportunities. We discussed common mistakes in grant applications, development of research questions for grants and manuscripts, career development of transdisciplinary expertise, areas of training that are fundamentally important for trainees to pursue MPE research, and tips for successful collaborations across disciplines.

30 May 2018

Session 1: Integrated genetic, epidemiologic, and tumor analyses #1 (Session Chair: Dr. Peter Campbell)

The first speaker of the meeting was Dr. Lorelei Mucci who presented an overview of her group’s work on integrating tissue biomarkers into prostate cancer epidemiology studies. She focused on two common molecular subtypes: the androgen-regulated gene fusion TMPRSS2:ERG and loss of the tumor suppressor PTEN. This work leverages prostate tissue biorepositories nested within the Physicians’ Health Study (PHS) and Health Professionals Follow-up Study (HPFS). The first set of studies focused on the high heritability of prostate cancer. Inherited variation within the androgen receptor (AR), which regulates AR expression, was associated with ERG-positive cancer but not ERG-negative disease [9]. Of 39 inherited prostate cancer risk loci, ten were differentially associated with risk when stratified by ERG status [10]. ERG-positive cancers show higher expression of the insulin and IGF1 receptors [11]. In unpublished data, vigorous physical activity was associated with a significantly lower risk of ERG-positive disease, whereas there was no association with ERG-negative disease. Finally, use of the cholesterol lowering drug, statins, was associated with a substantially lower risk of tumors showing PTEN loss. These data highlight the etiologic heterogeneity of prostate cancer, and the opportunities to elucidate discoveries based on integrating tissue biomarkers.

Dr. Nilanjan Chatterjee lectured on the development and utility of a broad mixed-effect two-stage logistic regression model for discovering new breast cancer germline genetic risk loci in the context of tumor heterogeneity [12, 13]. Breast cancers are highly heterogeneous, and it may become quickly inefficient to evaluate each potential risk locus with each different disease sub-type. With genome-wide association study (GWAS) data from the Breast Cancer Association Consortium (BCAC) that includes nearly 100,000 controls and a little over 100,000 cases, approximately 180 SNPs were discovered for breast cancer risk overall. When extending this work to subtype-specific analyses, a two-stage logistic regression approach was preferred over standard analyses to account for the large number of comparisons, correlations between markers, missing marker data, and other reasons. Case-control and case-case odds ratios were calculated using the mixed-effect two-stage approach, and 11 novel SNPs for subtype-specific breast cancers were identified. One of the more interesting findings from this work was the discovery that a TP53 SNP was associated with increased risk of luminal breast cancer and with decreased risk of triple negative tumors, probably reflecting the different pathologic mechanisms that drive these different tumor sub-types and potentially underscoring the broad utility of this statistical approach.

In the last lecture of the first session, Dr. Melissa Bondy spoke about GWAS results for gliomas, overall and when stratified by histological and molecular subtypes [14]. Brain tumors, which comprise a highly heterogenous group of cancers, account for 1–2% of all cancers overall. To better understand germline genetic risk factors for glioma, Dr. Bondy and collaborators pooled data from eight independent glioma GWAS datasets. Their combined meta-analysis identified 13 novel glioma risk loci (five for glioblastoma multiforme (GBM) and eight for non-GBM). Curiously, all but one locus showed significant allele frequency differences between GBM and non-GBM tumors. The only locus consistently associated with glioma risk was at 17p13.1 (TP53). Overall, genetic heritability is estimated to account for approximately 1/3 of the population variability in glioma. Further work showed a high degree of concordance, according to germline genetics, between familial and sporadic gliomas, suggesting genetic predisposition is largely the same for both types of disease. Many of the risk SNPs were in DNA-repair or telomere maintenance-related pathways. Future work in this area will include additional, detailed work within core pathways, sequencing efforts, and gene-environment interaction.

Session 2: Integrated genetic, epidemiologic, and tumor analyses #2 (Session Chair: Dr. Kana Wu)

The second morning session continued with the earlier theme of integrated studies in cancer epidemiology. Dr. Montserrat ‘Montse’ Garcia-Closas’s lecture gave an overview of recent developments in risk factor identification for breast cancer according to clinical subtype and the implications for those results on personal risk prediction. Breast cancer is a heterogeneous disease with different survival outcomes. The most aggressive tumors are hormone receptor negative tumors, including triple negative tumors that
are also negative for ERBB2 (HER2) amplification. These phenotypes represent approximately 13% of tumors and are more common in younger women and women of African descent. GWAS have identified nearly 180 germline loci for breast cancer risk overall. More recent work has identified clusters of SNPs associated with specific tumor features, including ESR-1 status and grade. Dr. Garcia-Closas and colleagues have developed Polygenic Risk Scores (PRS) for ESR-1+ and ESR-1-negative specific disease and are integrating them into risk models to identify women at different risks of breast cancer overall, and by subtypes; however, more work on implementation is needed before translating these tools into clinical practice [12, 13, 15–17].

Dr. Rulla Tamimi described the role of early life risk factors on markers in non-tumor tissues. Many risk factors for breast cancer in adults occur early in life. More specifically, there are windows of susceptibility that may be relevant to specific tissue markers of disease processes. Her group has conducted work on early life body size in the Nurses’ Health Studies (NHS) that asked women about their body size at ages 5, 10, and 20. They found that larger body size in early life was inversely associated with proliferative benign breast disease and breast cancer risk [18–20]. In a study of normal breast tissue adjacent to benign breast disease lesions, they found women who reported as being heavier early in life (ages 5–10) had a reduction of MKI67 (Ki67) expression [21]. In additional work, her group has examined breast tissue gene expression in the Nurses’ Health Studies using a transcriptome array. In preliminary work, they have seen differences in gene expression patterns related to exposures when considering tumor and adjacent normal tissue. Dr. Tamimi echoed a theme that was common during this year’s meeting: the need for more consortia work for validation, replication, and new discovery.

Dr. Christine Ambrosone discussed her research focused on understanding the etiology of more aggressive breast cancers in African American women, particularly tumors that do not express ESR1 (estrogen receptor, ER) that are associated with poorer prognosis [22]. She described results from a study of DNA methylation in tumors from African-American and European-American women that found that methylation of a gene important to guiding the luminal phenotype, FOXA1, was greatest in ESR1 (ER)-negative breast cancers [23]. Methylation was more common in women who had children and did not breastfeed, suggesting a mechanism for the increased risk of ESR1 (ER)-negative breast cancer with parity, and not breastfeeding [24].

Session 3: Integrated genetic, epidemiologic, and tumor analyses #3 (Session Chair: Dr. Song Yao)

After a 2-h pause in podium presentations for attendees to interact and to view poster presentations, the afternoon sessions began with the third installment of the integrated epidemiology theme.

The first lecture of the afternoon session was presented by Dr. Matty Weijenberg who discussed the role of MPE in supporting lifestyle guidelines for cancer prevention. She illustrated how MPE could contribute to the research directions provided by the third expert report from the World Cancer Research Fund (WCRF)/American Institute for Cancer Research (AICR) on diet, nutrition, physical activity, and cancer. First, she showed examples of how mutational spectra of tumors can provide clues to mechanisms, for example how heme iron intake is associated with specific G to A mutations in KRAS genes in colorectal tumors, pointing to the role of alkylating agents [25]. More recent studies suggest that tumor signatures of mutational processes are associated with exogenous mutational processes [26]. Second, she showed how exposure to the Dutch Hunger Winter was associated with a reduced risk of colorectal cancer only in tumors with a CpG-island methylator phenotype (CIMP) [27] and tumors with increasing number of IGFBP genes methylated [28]. Third, she showed how investigating subtypes of tumors can reveal previously unknown etiologies. For example, results from a meta-analysis revealed how adherence to a Mediterranean diet is associated specifically with a reduced risk of ESR1 (ER)-negative and PGR (PR)-negative postmenopausal breast cancer [29].

In the final lecture of the ‘integrated epidemiology’ sessions, Dr. Timothy Rebbeck discussed the role of germline genetics in explaining some disparities for prostate cancer [30, 31]. Prostate cancer has higher incidence and mortality rates in African American men compared to all other race/ethnic groups. Many prostate cancer susceptibility loci have been identified via GWAS for prostate cancer overall. But the contribution of these loci to prostate cancer disparities is unclear. To address this issue, Dr. Rebbeck’s group evaluated the population structure of 68 previously identified prostate cancer susceptibility loci by calculating: (1) genetic disparity contribution statistics to quantify the contribution of each SNP to differences in prostate cancer risk across populations, and (2) genetic risk scores that integrate GWAS results with allele frequency data from 45 African and 19 non-African populations. They found that predicted prostate cancer risks were highest for men of West African descent and lowest for men of East Asian descent. These population-level differences were further explained by the out-of-Africa bottleneck and natural selection. Only a few loci seemed to drive the excess prostate cancer risk observed in African American men. Although most prostate cancer
susceptibility loci are evolving neutrally across different race/ethnic groups, there are several instances where alleles have hitchhiked at higher frequencies with adaptive alleles, including alleles for skin pigmentation (at 2q37).

**Session 4: Immunology, immunotherapy, and prevention #1 (Session Chair: Dr. Amanda Phipps)**

The first lecture in the ‘immunology-MPE’ session was given by Dr. Shuji Ogino who spoke about the broad importance of incorporating immune system data into MPE studies. Despite remarkable advances of cancer immunology in recent years, investigations on the influences of the exposome on tumor-immune interactions lag. To address a substantial gap between cancer immunology and epidemiology, the integrative field of immunology-MPE can investigate influences of the exposome (dietary, lifestyle, environmental, microbial, pharmacological, and other exposures) on tumor-immune interactions [32, 33]. Using epidemiological studies and colorectal cancer cases with data on immune response, tumor molecular pathology, and tissue microenvironments, proof-of-principle immunology-MPE studies provide evidence supporting hypotheses that several exposures influence carcinogenic processes through their influences on tumor-immune interactions [34–38]. For instance, marine omega-3 polyunsaturated fatty acid intake has been associated with a lower risk of colorectal carcinoma containing abundant FOXP3+ cells (mostly regulatory T cells) [34] and a prudent dietary pattern has been associated with a lower risk of colorectal carcinoma containing abundant *Fusobacterium nucleatum* [38]. These new insights from immunology-MPE research can provide a possible path for precision immunoprevention and immunotherapy.

Dr. X. Shirley Liu introduced her work in mining and integrating large-scale tumor molecular profiles to inform cancer immunology and immunotherapy. Dr. Liu discussed three algorithms that her laboratory developed to extract useful insights from treatment-naïve RNA-seq samples in The Cancer Genome Atlas (TCGA). First, ‘TIMER’ can estimate immune cell components in tumors [39], and a webserver was created for users to explore immune infiltration across TCGA tumors and to draw inferences on user-provided samples [40]. Second, ‘TRUST’ can assemble T cell receptor (TCR) and B cell receptor (BCR) complementarity-determining regions (CDR3s) from bulk tumor RNA-seq data [41]. When applied to over 10,000 samples in the TCGA, TRUST assembled 3 M TCR CDR3 sequences from tumor RNA-seq samples and revealed associations between tumor infiltrating TCR clonotype diversity and tumor mutational load [42]. TRUST also identified 30 M BCR sequences from TCGA tumor RNA-seq and revealed widespread B cell clonal expansions among other events [43]. Third, ‘TIDE’ software derived gene expression signatures from pretreated tumor specimens to predict patient response to anti-PDCD1 (PD-1) and anti-CTLA4 treatment [44]. TIDE analyses of published immune checkpoint inhibitor trials suggested some tumors are unlikely to respond to anti-PDCD1 (PD-1) or anti-CTLA4 alone. This work indicates that tumor RNA-seq, even on treatment naïve tumors, is cost-effective to inform tumor microenvironment and immunity.

Dr. Robert ‘Rocky’ Schoen described his group’s experiences in developing a vaccine for colorectal cancer. Immunotherapy targeting of antigens that are aberrantly expressed on colon cancers and polyps offers the potential for relatively non-invasive, non-toxic, and prolonged preventive strategies. Whereas vaccines in advanced cancers have had little success, likely because of immunosuppressive tumor microenvironments, vaccines administered in pre-malignant stages when the immune system is still powerful should be more effective. Dr. Schoen and colleagues are testing this hypothesis using the adenomatous polyp-to-colon-cancer pathway. Their target is MUC1, a tumor-associated antigen that is abnormally expressed on polyps and colon cancers. In a pilot study of 39 patients with a history of advanced adenomas, after a series of 3 injections, almost half of the patients showed a twofold ratio increase in anti-MUC1 IgG at week 12 compared to pre-vaccination levels. A booster injection at week 52 resulted in a large increase in IgG, demonstrating a persistent T cell memory response [45]. There was minimal evidence of toxicity. A double-blind randomized trial of 110 patients using a similar vaccination protocol is due to report in late 2019. That trial will also evaluate a clinical endpoint, via assessment of adenoma recurrence.

**Session 5: Immunology, immunotherapy, and prevention #2 (Session Chair: Dr. Xuehong Zhang)**

In the final session of the day, Dr. Catherine Wu lectured on her work in identifying tumor antigens. Multiple lines of evidence demonstrate that tumor neoantigens are an important class of immunogenic antigens. Neoantigens arise from amino acid changes encoded by somatic mutations in the tumor cell. This work has advanced in recent years due to the availability of next-generation sequencing approaches and the maturation of predictive algorithms [46]. One of the central questions of Dr. Wu’s work is: can a personalized cancer vaccine stimulate anti-tumor immunity in humans? Her group conducted a trial in high-risk melanoma patients. They dosed 6 melanoma patients with up to 20 neoantigens to test for safety, feasibility, and immune response. Across the 6 patients, they observed ~20% CD8 and >60% CD4 T cell responses against the neoantigens [47], all of which were new responses following vaccination. In another trial, Dr. Wu and colleagues tested Neovax in patients with...
In the final presentation of the day, Dr. Marios Giannakis presented work on investigating the genomic mechanisms of immune evasion in colorectal cancer. Immune checkpoint blockade has shown activity in approximately 50% of MSI-high colorectal cancers while it is ineffective in microsatellite stable tumors [50, 51]. To better understand the genetic drivers of immune evasion in colorectal cancer, Dr. Giannakis and colleagues integrated next generation sequencing data from over 1,200 tumors with transcriptional and immunohistochemical measures of immune infiltration. The tumor samples for their studies were from TCGA, the NHS, and the HPFS [52]. They demonstrated that WNT-signaling and immune-related genes were significantly mutated in colorectal cancer. They also found frequent inactivating antigen-presentation machinery mutations in MSI-high tumors, and an inverse association between WNT-signaling activity and T cell infiltration in all subtypes of colorectal cancer. Tumors with biallelic disruptive mutations in APC or with AXIN2 super-enhancer hypomethylation had a significantly decreased T cell transcriptional signature [53]. In summary, this work found evidence of immuno-editing through disruptive mutations in antigen-presentation machinery, and of exclusion of an effective immune response through an active WNT-signaling pathway in colorectal cancer. These results shed light to the underlying molecular mechanisms of immune evasion in this disease.

1 June 2018

The second day of the meeting began with award announcements to trainees and early career investigators.

Session 6: Novel disease phenotyping in future medicine and population science #1 (Session Chair: Dr. Jonathan Nowak)

The first lecture of the day was given by Dr. Jeffrey Golden who first shared a few clinical case reports whereby modern tools in molecular pathology aided in more precise diagnoses and superior treatments for patients and almost certainly improved their prognoses. Dr. Golden summarized the importance of computational pathology [54, 55] into five central components: [1] mutation-specific treatment stratification, including clinical trial eligibility; [2] better precision diagnostics; [3] superior prognostication; [4] better targeted therapies, which may be more effective, less toxic, and more cost-effective; and [5] new biomarker discovery. One of the central challenges for a practicing clinician is in translating the abundance of computational data generated by omics platforms (e.g., Oncopanel) into tractable information and, ultimately, into knowledge of what those mutations mean for the benefit of the patient. Dr. Golden’s institution has converted these sorts of information into knowledge—that is, what mutations in those genes mean to the patient and clinician [56]. He used Google maps’ system of layered geospatial databases as an analogy for creating relational databases for patients, based on a multitude of patient (e.g., clinical, omic, pharmacy) and external (e.g., electronic medical records, published literature) inputs, that are ultimately led back to actionable decisions for clinicians and their patients.

Dr. Lynette Sholl’s lecture focused on recent advancements in liquid biopsy techniques for solid tumors. She started with a clinical case report that highlighted some of the advantages of liquid biopsies over tumor/solid tissues, including: tumor tissues often have limited mass and/or normal cell contamination; there is often a need to repeat a solid tissue biopsy upon relapse; and some tumors are anatomically inaccessible. Liquid biopsies can look for circulating tumor DNA (ctDNA) although their rarity in peripheral blood makes the approach challenging. Current and emerging technologies include commercial allele specific PCR platforms, droplet digital PCR (ddPCR), NGS, and electric field-based measurements [57]. Investigators at her institute decided to invest in ddPCR for ctDNA testing in cancer patients. Using ddPCR for EGFR and KRAS hotspots, they found sensitivity was a direct correlate of the number of metastatic sites, owing to the amount of DNA shed into circulation. Patients with only 1 metastatic site saw sensitivity levels of approximately 60% whereas patients with four or more metastatic sites had nearly 100% sensitivity with ddPCR [58]. Ultimately, liquid biopsy may be used in detection of early relapse after definitive therapy, minimal residual disease testing, or even cancer screening.

Dr. Yujing (Jan) Heng’s talk highlighted the importance of gene expression pre-processing methods to obtain reliable and reproducible breast cancer molecular subtype classification by PAM50. Her work compared two established pre-processing methods (i.e., modified median gene centering [59] and subgroup-specific gene centering [60]) to compute molecular subtypes using PAM50 in tumor and tumor-adjacent tissues from participants in the NHS. She reported that although molecular subtypes were highly comparable using either method, the subgroup-specific method tends to classify more cases into more aggressive subtypes. The distribution of molecular subtypes within the NHS/NHS-II was comparable to other population-based studies, and as expected, there were more cancer recurrences in women with Basal-like subtype compared to Luminal A subtype.
Lastly, she showed that the correlation of molecular subtypes classified using PAM50 and immunohistochemical (IHC) surrogates remain poor and more research is needed to refine the IHC definitions to more closely approximate PAM50 subtypes [61].

**Session 7: Novel disease phenotyping in future medicine and population science #2 (Session Chair: Dr. Reiko Nishihara)**

After a short break, the second session of the day continued with the ‘novel disease phenotyping’ theme. The first lecture of the session was given by Dr. John Quackenbush who described his group’s efforts to infer biological networks from GWAS and gene expression data. To do this, they developed several systems biology algorithms [62, 63]. Dr. Quackenbush’s lecture described how these algorithms were used to better understand networks of GWAS-identified SNPs and gene expression data with outcomes, and how patterns of gene networks differ between phenotypes. Central to this work are the hypotheses that biological systems are driven by complex networks; the structure of the network captures the biology of the system; and, that network structure is conditional, depending on tissue, biological state, and individual. One of the main findings from their work is that biological networks are organized into tight communities such that, in most instances, it is not a single gene controlling a single trait, but a family of genetic variants that influence a process. They also find disease-associated (i.e., GWAS) SNPs map to communities whose genes share functions that are related to the disease, and that most GWAS SNPs are not global hubs in the network, but local hubs in the network. Overall, their work demonstrates how network analysis can take us beyond simple differential expression in understanding disease [64, 65].

Dr. Hugo Aerts described his work at the intersection of radiology, bioinformatics, and data science. He discussed recent work of building Artificial Intelligence (AI) image analysis systems to extract a rich radiomics set and used these features to build biomarkers [66]. He illustrated how technological advances in AI and deep learning are moving imaging modalities into the heart of patient care as imaging can address a critical barrier in precision medicine because solid tumors can be spatially and temporally heterogeneous, and the standard approach to tumor sampling, often invasive needle biopsy, is unable to fully capture the spatial state of the tumor [67, 68]. The main objectives of the talk were to learn about the motivation and methodology of AI technologies in radiology, to learn about the existing and future potential role of radiologic AI with other omics data for precision medicine, and to learn about open-source informatics developments [69].

**Session 8: Discussion topic special lectures (Session Chair: Dr. John Quackenbush)**

In the final session for podium presentations, the organizing committee selected five submitted abstracts whose topics were thought to provide the broadest interest and potential to generate discussion for the attendees.

Dr. N Sertac Kip lectured on her work in developing a liquid biopsy for tumor detection. Dr. Kip spoke of the limitations to standard tumor profiling (e.g., biopsy/resection), including invasiveness, pain, cost, and lack of sensitivity to mutations unique to the non-biopsied tissue [70]. Further, targeted therapies place tumor cells under selective pressure, thereby triggering clonal progression, which can then be captured [71]. Liquid biopsy is ideal for comparing pre- and post-treatment variants and to optimize sequence of therapy [72]. Lung cancer appears to be an ideal tumor site for this application. Liquid biopsy measures biomarkers that are predictive and prognostic for lung cancer [73]. Coming back to the title of her talk, “How solid is liquid biopsy?”, only two of 34 cases had discordance between solid tumor and liquid biopsy measures, indicating that the liquid biopsy is quite solid [74]. There are some people with mutations in tissue that are not detected in liquid biopsy. Sensitivity also varies by tumor type and stage [75]. Research is rapidly evolving in this area, but the techniques are not yet suitable for wide clinical application and guidelines are currently lacking for standardization of the liquid biopsy results.

Dr. Camila Lopes-Ramos spoke about her work in identifying biological explanations for the higher incidence and mortality rates for men than women from colon cancer. Lifestyle and serological (i.e., sex steroid hormone) differences are often postulated to explain some, but not all, of these observed sex differences. The potential molecular features that drive sex differences are understudied. Her work used both transcript-based and gene regulatory network methods to analyze RNA-seq data from TCGA for colon cancer. They found no meaningful differences between tumors from men and women for gene expression. Next, they examined patient-specific gene regulatory networks and found considerable sex differences in drug and xenobiotic metabolism via cytochrome P450 pathways which were considerably more pronounced in women. This finding was replicated in several independent study samples. This drug metabolism pathway was not associated with survival in men; however, women treated with chemotherapies that had increased targeting (compared to less targeting) of this pathway had considerably better 10-year overall survival [76]. This network-based approach can be applied to explore other etiologic and demographic differences for cancer and other complex diseases.

Dr. Molin Wang lectured on the problem of sample selection bias due to tissue availability and solutions to this problem. Missing data is a common problem for tumor subtype
data. This poses severe statistical challenges for MPE research because the outcome is missing. She reported the percentages of colorectal, breast and ovarian cancer cases with missing tumor marker data in representative national and international cohort studies and consortia. She introduced the definitions and assumptions of possible missingness patterns of the tumor marker data in the MPE context and used the NHS breast and colorectal cancer data to illustrate the problem of sample selection bias in MPE research. She then described the statistical methods available to explore these issues, such as the complete case analysis method, the missing indicator method, inverse probability weighting [77], and the multiple imputation method. They found that the former two methods could lead to biased estimates and the latter two methods were usually most helpful in dealing with the potential selection bias problem. She used a colorectal cancer molecular subtype to illustrate these methods. She described the possible scenarios when the missingsness is not at random, and the statistical challenges in these scenarios.

Dr. Song Yao presented recent data from his group on population differences between women of African American descent and European American descent in the breast tumor immune microenvironment. Convincing evidence demonstrates marked differences in systemic immune response between African American and European American populations [78–80], which is also supported in a recent study on circulating cytokine levels [81]. However, data are scarce on population differences in tumor immune microenvironment. With data from the Women’s Circle of Health Study (WCHS), they showed that breast tumors from African American women had a significantly stronger presence of tumor-infiltrating lymphocytes than breast tumors from European American women, independent of tumor histopathological features. Using NanoString immune profiling, they confirmed the overall stronger immune infiltration in breast tumors from African American women than those from European American women, and further showed stronger exhausted T cell signatures in tumors from African American women. Their data revealed marked population differences in tumor immune response, which may contribute to some of the observed racial disparities in breast cancer survival.

Dr. Kun-Hsing Yu lectured about his work on integrating lung cancer multi-omics and histopathology images [82]. Lung cancer is the most prevalent cancer worldwide, and histopathological assessment is indispensable for its diagnosis [83]. However, how histopathology findings relate to molecular abnormalities remains largely unknown, and human evaluation of pathology slides do not accurately predict prognosis. To address this gap, his group obtained over 2,100 hematoxylin and eosin stained histopathology whole-slide images, RNA sequencing, and proteomics data of lung adenocarcinoma and squamous cell carcinoma patients from TCGA, and nearly 300 additional images from the Stanford Tissue Microarray (TMA) Database [84]. They extracted nearly 10,000 quantitative image features and used regularized machine-learning methods to select the top features and to distinguish shorter-term survivors from longer-term survivors with stage I adenocarcinoma or squamous cell carcinoma in the TCGA data set [85]. They successfully validated the survival prediction framework with the TMA cohort [85], identified the cell-cycle regulation and nucleotide binding pathways underpinning tumor cell dedifferentiation [86], and built an integrative histopathology-transcriptomics model to generate better prognostic predictions for stage I adenocarcinoma patients compared with gene expression or histopathology studies alone [86]. These results suggest that automatically derived image features can predict the prognosis of lung cancer patients and therefore contribute to precision oncology.

Conclusions

The Fourth International MPE Meeting assembled over 170 trainees and experts working in the various, diverse scientific disciplines that comprise MPE. As even more sophisticated means of molecular characterization of disease processes enter epidemiologic studies and clinical medicine, the utility and preponderance of MPE principles and methods should continue to expand. As actively discussed in the Fourth International MPE Meeting, new ideas of flexibly shaping and integrating multiple disciplines are further expanding opportunities in biomedical and population sciences [87]. In terms of key recommendations and next steps, integration of tumor immunology into epidemiologic studies and further exploration of disparity research were concluded as high priorities for the field. Additionally, further collaborations between computational and population-level scientists were noted as high priority as were general pooling projects between studies with similar data. We look forward to meeting again at the Fifth International MPE Meeting, tentatively planned for June 2020 in Boston, MA, USA.

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