Germline genomes have a dominant-heritable contribution to cancer immune evasion and immunotherapy response

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Background: Immune evasion is a fundamental hallmark for cancer. At the early stages of tumor development, immune evasion strategies must be implemented by tumors to prevent attacks from the host immune systems. Blocking tumors’ immune evasion will re-activate the host immune systems to eliminate tumors. Immune-checkpoint therapy (ICT) which applies anti-PD-1/PD-L1 or anti-CTLA4 treatment has been a remarkable success in the past few years. However, ~70% of patients cannot gain any clinical benefits from ICT treatment due to the tumor-immunity system’s complexity. In the past, germline pathogenic variants have been thought to have only minor-heritable contributions to cancer.

Results: Emerging evidence has shown that germline genomes play a dominant-heritable contribution to cancer via encoding the host immune system. The functional components of the immune system are encoded by the host genome, thus the germline genome might have a profound impact on cancer immune evasion and immunotherapy response. Indeed, recent studies showed that germline pathogenic variants can influence immune capacity in cancer patients at a population level by (i) shaping tumor somatic mutations, altering methylation patterns and antigen-presentation capacity or (ii) influencing NK cell’s function to modulate lymphocyte infiltration in the tumor microenvironment. In addition, the HLA (types A, B or C) genotypes also shape the landscape of tumor somatic mutations.

Conclusion: These results highlight the indispensable roles of germline genome in immunity and cancer development and suggest that germline genomics should be integrated into the research field of cancer biology and cancer immunotherapy.

Keywords: germline; genomics; cancer; immune evasion; immunotherapy response

Author summary: Traditionally, it has been believed that germline pathogenic variants and family histories explain 5%–10% of cancer patient population, thus, heredity has been suggested to have a small contribution to tumorigenesis and metastasis. However, the host immune system often interacts with cancer cells, therefore, escaping from the host immune system surveillance is one of the critical means for tumorigenesis. In the past a few years, it has been shown that germline pathogenic variants influence immune capacity in cancer patients at a population level. From the cancer-immune system point view, heredity plays a dominant role in tumorigenesis, metastasis and immunotherapy response.

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INTRODUCTION

Immune evasion is a fundamental hallmark for cancer cell survival

One of the cancer hallmarks is immune evasion [1]. During the initiation and development of tumors, the host immune system monitors “foreign” cells such as cancerous cells in the human body. Generally, the immune system can eliminate these mutated cells, however, these cells can find ways to escape from the host immune system by developing various immune evasion strategies. For example, tumor cells can deregulate pathways such as antigen presentation [2] and interferon-gamma (IFN-γ) pathways [3] or activate intrinsic signaling pathways such as mitogen-activated protein kinase (MAPK) [4,5] and Wnt signaling pathways [6,7] to block tumor-infiltrating lymphocytes (TILs) from entering the tumor immune microenvironment (TIME). Without implementing these immune evasion strategies, a tumor cell can be quickly removed by the host immune system.

Immunotherapy represents a truly exciting therapeutic success in oncology. Particularly, the discovery of immune “brakes” or checkpoint proteins, such as cytotoxic T-lymphocyte-associated protein 4 (CTLA4), programmed cell death protein-1 (PD-1) and programmed cell death protein ligand-1 (PD-L1), has led to developing various strategies to overcome these immune checkpoints which could further remove the immune “brakes”, allowing T cell activation and elimination of tumors. In the past few years, ICT has shown an overall response rate of 10%–40% in patients with melanomas and other types of cancer [8,9]. However, for the rest of these patients, ICT failed to show their intended effect.

ICT response is dependent on many factors. For example, tumor antigens need to be released and presented onto the tumors’ cell surface so that immune cells such as T cells will be activated and infiltrated into the tumor microenvironment. Further, anti-PD-1/PD-L1 or anti-CTLA4 treatment helps in overcoming an immunosuppressive microenvironment for cytotoxic T lymphocytes (CTLs) to recognize and ultimately kill these tumor cells [10]. TIME interacts with cancer cells to influence metastasis and clinical outcomes [11,12]. Therefore, a deeper understanding of TIMEs and identification of key germline regulators modulating TIMEs and immune-evasion strategies would shed further insights into ICT response, resistance and might improve existing immunotherapies. For example, it has been known that rich TILs are critical for ICT response [13], and thus understanding how TILs are affected by germline genomics will provide a new avenue for ICT treatment. Due to the relatively short period of a wide application of immunotherapy, the roles of germline genomics in cancer immunity and immunotherapy have not been extensively studied. Nevertheless, several pilot studies have been conducted in this direction. In this review, we will discuss the recent progress and its implications for future research in this area.

Germline pathogenic variants have been shown to have only minor-heritable contributions in cancer in the past

Traditionally, inherited germline variants have been shown to account for a relatively small fraction of cancer patients. For example, the most famous BRCA1/2 germline pathogenic variants account for 5%–10% of breast or ovarian cancer [14,15]. The effect sizes of other germline variants are relatively small, and thereby have a minor influence on cancer development [16]. Besides, many germline variants are haplotypes and thus do not have a significant impact on cancer development as well. Albert Knudson was the first scientist who discovered germline variants in 1971. He studied patients with retinoblastoma carrying mutations in RB1. At that time, it was already known that two alleles existed for each gene, and a mutation in both copies of RB1 would be required to develop retinoblastoma. Knudson noticed that bilateral retinoblastomas (i.e., retinoblastoma in both eyes) were more frequent in younger patients (25%–30% of all cases), suggesting that germline pathogenic variants were involved. As mutations in both alleles of RB1 are required to develop retinoblastoma, inherited germline mutation in this gene was predisposing the children to cancer. Knudson’s findings were later described as the “two-hit” hypothesis, supposing that children possessing an inherited RB1 mutation need to obtain a second mutation to develop retinoblastoma [17,18]. In contrast, children without RB1 pre-disposition were less likely to develop retinoblastoma as they needed to acquire two mutations. Mice with germline inactivation in Rb1 and p107 (another Rb family member) developed unilateral retinoblastoma by nine months [19], while with germline deficiency in Rb1 and p130 (another Rb family member) developed bilateral retinoblastomas with a 100% of penetrance in four months [20]. Today, multiple hereditary germline mutations have been associated with cancer. For example, people carrying a dysfunctional germline p53 gene have a ~90% chance of developing a variety of cancers before the age of 70 [21]. Similarly, mice deficient of germline p53 are extremely susceptible to radiation-induced tumorigenesis [22,23]. p53 is widely regarded as the most important tumor suppressor in the human genome. It acts as one of the major gatekeepers in cancer and has multiple functions such as ensuring genomic integrity and...
promoting DNA repair, cell cycle regulation and apoptosis, signal transduction and cell adhesion [24]. Other cancer-related inherited germline pathogenic variants include those found in BRCA1/2, which are tumor suppressor genes involved mainly in DNA repair and are associated with breast and ovarian cancer [25,26]. In addition, germline pre-dispositions in BRCA1/2 in women have been associated with an increased risk of developing breast and ovarian cancer by the age of 80 (69%–72% and 17%–44%, respectively) [27]. Although the prevalence of BRCA1/2 germline pathogenic variants worldwide is very low (5%–10%), they are now regularly checked in genetic screening. Other heritable genes include PTEN, whose germline mutations induce Cowden syndrome [28], and APC, which has been associated with familial adenomatous polyposis [29] and is accurately modeled by germline mutations in mouse models [30,31]. RET and MEN1 germline mutations have been shown to cause endocrine cancers while VHL alterations result in kidney and other types of cancer [32]. Moreover, Lynch syndrome, a form of colorectal cancer, is linked to germline mutations in MSH2, MLH1, MSH6, PMS2 and EPCAM [33]. A lot of these germline mutations could be or have been modeled by experimental approaches such as using CRISPR/Cas9-modified mice [34,35]. Investigation of whole exomes in cancer patients revealed that the germline genomic landscape of these patients is informative and can provide insights on tumor development [36,37]. However, these germline variants have relatively low penetrance and can only explain the heredity for 5%–10% of cancer patients [37,38]. Therefore, germline genomic information has been largely ignored in cancer research due to its limited role in cancer development in the past.

**GERMLINE PATHOGENIC VARIANTS MODULATING CANCER IMMUNE EVASION AND IMMUNOTHERAPY RESPONSE ARE HERITABLE ACROSS THE WHOLE CANCER PATIENT POPULATION**

As shown above, germline pathogenic variants are thought to have only minor contributions to cancer, and thus have been largely ignored in cancer research. As systems biology emerges and develops in the past 10 years, it is now possible to identify germline variants which encode proteins in meaningful processes such as the immune system that could have a heritable influence on cancer development. In the cancer hallmark network framework [39], we proposed the logics of complex cancer biology as a myriad of phenotypic complexities governed by a limited set of underlying organizing principles. Thus, we expected that germline pathogenic variants in some of these organizing principles (or cancer hallmarks) could have a heritable influence across the whole cancer patient population, and thus play a dominant-heritable role in cancer development and metastasis. In particular, in the cancer-immunity system, the immune components are not so heterogeneous as tumor cells are, therefore, germline pathogenic variants related to the immune system could have a heritable influence on immune evasion and immunotherapy dominantly. The role of germline pathogenic variants on the cancer-immunity system has been rarely investigated. Recently, we and others pursued this direction using omics data collected from The Cancer Genome Atlas (TCGA) and showed that germline pathogenic variants influence cancer-immunity and immunotherapeutic response via different mechanisms (Fig. 1).

**Germline genomes provide evolutionary and immune constraints to tumor somatic mutations across the whole cancer patient population**

Although cancer somatic mutations seem chaotic, complex and unorganized, they have been selected by the host-specific genetic makeup to become the components of cancer cell machinery representing cancer hallmarks [39]. In the cancer hallmark network framework [39], we have proposed that the pre-existing genetic makeup could restrain somatic mutations and cancer clonal evolution to form specific signaling pathways and regulatory networks to represent cancer hallmarks. Thus, the cancer hallmark network framework suggests that based on host genome information, a cancer patient’s somatic mutations in his/her tumors are predictable, and therefore, the tumor recurrence and metastasis are also predictable.

To demonstrate these concepts, we have developed a novel network-based computational algorithm, eTumorMonitor, which predicts key somatic mutating genes based on host genomic information (Tibiche et al., unpublished data). By applying this algorithm to the germline whole-exome genomic data of breast, lung, and prostate cancer patients, we were able to predict 20–30 key somatic mutating genes in the cancer founding clone, which is the first cancer cell inferred from tumor sequencing data, for each patient. The predicted somatic mutating genes were enriched 10,000 times comparing to random groups. The prediction applies to all the cancer patients tested (n > 600, Fig. 1). These results support the notion in the cancer hallmark network framework that host pathogenic variants play an important role in selecting somatic mutating genes during cancer clonal evolution. These key somatic mutating genes in tumor cells will be organized into several tumor cell-intrinsic
signaling pathways to maintain the malignant growth of cancer. Key tumor cell-intrinsic signaling pathways such as MAPK, Wnt and TGF-β pathways are associated with cell proliferation and metastasis [40]. Also, recent works showed that blocking TILs in TIMEs is an intrinsic feature for germline pathogenic variants [41–43]. Thus, host germline genomes could regulate TIMEs and immunotherapy response by selecting key cancer somatic mutating genes, pathways and regulatory networks. Rethinking of the constraints on cancer somatic mutations by host-specific genetic makeup would provide a novel window to explore and understand the complexity of tumor-host interactions.

Except for the functional constraints exerted on cancer somatic mutations by germline genomes, as shown above, HLA genotypes can exert an immune constraint on the landscape of tumor somatic mutations as well (Fig. 1). Major histocompatibility complex (MHC) class I (HLA-A, -B and -C) is an important machinery for antigen presentation of peptides for intracellular proteins including cancer-driving proteins onto the cell surface. CD8⁺ T cells can recognize “foreign antigens” which are derived from mutated proteins and thus kill the cancer cells. The binding affinity of the MHC-I complex for peptides is a major determinant of antigenicity: if a peptide can bind to the MHC-I with a higher binding affinity, it can be easier to be presented onto the cell surface so that T cell can recognize it. MHC-I binding affinity is dependent on the genotypes of three HLA genes (HLA-A, HLA-B and HLA-C) which are highly polymorphic. Thus far, more than 10,000 HLA alleles for the three genes have been documented [44]. The combination of six MHC-I alleles in each person determines which peptides can be more effectively presented. Thus, each person may have an MHC-I complex that possesses different binding affinity to a same peptide. If a mutated peptide can be effectively presented, the corresponding cell can be eliminated by T cells during the early stages of tumor development. A computational analysis of the somatic mutations/peptides in thousands of tumors has shown that MHC-I genotype influences the occurrence of specific oncogenic mutations, and recurrent oncogenic mutations are enriched in the peptides that are poorly presented by MHC-I complexes [45]. Somatic mutations encoding poorly
presented peptides were more frequently observed in tumors [45]. The restriction on the landscape of cancer somatic mutations in tumors by MHC-I genotypes is not limited in a subset of patients but actually in almost all patients we tested (i.e., several thousands of cancer patients collected in TCGA) [45]. These results suggest that germline HLA-A, HLA-B, and HLA-C variants restrict cancer somatic mutations via immunoediting during tumor formation. One interesting question remains: are people who have certain HLA genotypes having a higher risk of developing solid cancer?

It should be noted that one drawback of this study [45] is that they used all the somatic mutations in the tumors to examine the immune-selection pressures from the HLA phenotypes. Based on the timing of individual tumor evolution, their somatic mutations can be classified into two groups: founding clonal mutations and subclonal mutations. Genomic and epigenetic alterations in the process of transformation of a normal cell into a cancer founder cell in a tumor are called founding clonal mutations or alterations, while the rest are called subclonal mutations or alterations [39,46–48]. While there are a few different opinions [49,50], the theory that tumor originates from one founding clone is commonly accepted and validated by many studies [51–58]. We hypothesized that immune-escaping strategies exist before the formation of a founding clone for each tumor. As mentioned above, these strategies could include constraining somatic mutations through antigen presentation pathway (related to HLA gene polymorphism), deregulating TILs in TIMEs through INF-γ, MAPK and Wnt pathways, or altering expression of PD-1/PD-L1 and CTLA4 checkpoint proteins. Therefore, once a founding clone is formed, the immune-selection pressure on somatic mutations from the MHC-I system could be dramatically reduced or even disappeared, meaning that the immune-selection pressure on somatic mutations from the MHC-I system could be much less in subclones. Nevertheless, selection pressure from HLA genotypes could largely play its role in the early stages of cancer development, namely, before the formation of a founding clone. These immune evasion strategies may shape signaling pathways in cancer founding clones, which in turn, influence the signaling pathways in their subclones by functionally selecting somatic mutations to enhance cancer hallmarks [39].

**Germline genomes influence tumor recurrence via immune-related genes**

To further prove the correctness of these concepts proposed in the cancer hallmark network framework mentioned above, we have developed another computational algorithm, eTumorMetastasis [59], which transforms germline pathogenic variants into network-based profiles and identifies network operational gene signatures. These gene signatures model the tipping point at which a tumor cell shifts from a state that does not favor recurrences to one that does. In addition, to improve prediction accuracies and robustness, eTumorMetastasis incorporates key concepts of the multiple survival screening (MSS) algorithm [60,61] developed by our lab previously. By applying the eTumorMetastasis to whole-exome germline genomic sequencing data of ER + breast cancer patients [48], we showed that gene signatures derived from the genes encoding germline pathogenic variants significantly distinguished recurred and non-recurred patients in ER + breast cancer cohorts (n = 200 and 295, respectively, Fig. 1). Furthermore, the germline-based prediction significantly outperforms the well-known Oncotype DX test (i.e., Oncotype DX breast cancer recurrence test) which is widely used in clinics to help doctors making decisions for clinical treatments by determining the probability of recurrence based on the expression of 21 genes in the Oncotype DX 21-gene assay in breast tumors [62]. It should be noted that our predictions are based on the whole ER + breast cancer population, suggesting that host germline pathogenic variants could provide a profound constraint on the evolution of tumors and therefore exert a contingent effect on tumor metastasis and clinical outcomes. The performance of our algorithm (eTumorMetastasis) [48,59] strongly supports the notion proposed in the cancer hallmark network framework [39]. Therefore, we proposed that germline genomic information of cancer patients could predict cancer recurrence in a population-wide manner [59].

Practically, eTumorMetastasis provides a new possibility for developing blood tests to predict prognosis in different types of cancer. Traditionally, the prediction of tumor recurrence (such as the Oncotype DX assay) is based on sequencing data of tumor samples. To conduct the test using tumor samples, an invasive, often risky and painful procedure is used, either through a needle or surgery. However, our eTumorMetastasis-based test is utilizing “liquid biopsy”, which uses white cells from blood [48,59].

In addition, germline pathogenic variants are predominantly enriched in the processes related to T cell function, antigen presentation and cytokine interactions [43], suggesting that inherited germline variants are likely impairing the adaptive and innate immune systems in recurred samples. These results provide a new angle to understand the host-tumor interaction that germline pathogenic variants of immune-related genes could play an important role in metastasis.
Germline genome sequence patterns shape oncogenic pathways, cancer risk, and tumor recurrence

Germline genomes have a dominant-heritable role in cancer. To identify key regulators that deregulated chromatin landscape of tumors plays an important role in modulating the immune landscape and TIMEs, we classified ~6,000 tumors from TCGA representing 13 common cancer types into TIME-rich, -intermediate and -poor subgroups which contain high, intermediate or low levels of TILs, using tumor RNA-seq data and by clustering analysis [64]. Analysis of these three subtypes indicated that TIME-rich tumors have a significantly better prognosis than TIME-poor tumors. By applying the TIME subtypes to ICT clinical trial samples, we showed that anti-PD-1 responders were significantly enriched in TIME-rich tumors, while almost none were found in TIME-poor tumors [64]. Genome-wide analyses of germline variants in the 6,000 patients showed that inherited germline pathogenic variants in natural killer (NK) cells are significantly associated with the TIME-poor subtype [64]. The number of inherited germline pathogenic variants in NK cells are negatively associated with TILs’ abundance, metastasis, survival and ICT response, thus positively correlated with the risk of developing cancers [64]. These results suggest that germline variants in NK cell-related genes not only affect NK cells, but also affect other types of immune cells (such as T cells) from entering the TIMEs, and thus increase the risk of developing tumors. Notably, the conclusions are drawn from ~6,000 cancer patients representing 13 common cancer types, suggesting that germline pathogenic variants in NK-related genes play an important role in modulating TILs’ abundance in TIMEs, ICT response and clinical outcomes in a dominant manner (Fig. 1).

Germline pathogenic variants shape DNA methylation patterns and TIMEs in tumors

Other than genomes, the epigenome provides another layer of gene regulation on transcriptional programs, which impacts phenotypes and cellular memory. Epigenomic changes have been well-documented in cancer development [65]. Somatic mutations in genes such as chromatin-remodeling factors [66], histone-modifying enzymes [67], and DNA methylation related enzymes [68] have also been found in tumors. By comparing differentially expressed genes between TIME-rich and TIME-poor tumors in TCGA, we found that chromatin-remodeling related genes are significantly associated with TIME-poor tumors (unpublished data), suggesting that deregulated chromatin landscape of tumors plays an important role in modulating the immune landscape and TIMEs. Activated SWI/SNF chromatin remodeling complex in tumor cells inhibits the activation of IFN-pathway [69,70]. Inactivation of IFN-pathway to reduce IFN-γ production is one of the common immune evasion strategies for tumors [71,72]. Furthermore, the deletion of PBRM1, a subunit of the SWI/SNF chromatin remodeling complex, increased response rates to anti-PD-1/PD-L1 treatment in cell renal cell cancer patients [69,70]. These results suggest that the deregulation of DNA methylation programs plays a key role in the regulation of TIMEs and ICT response. It should be noted that epigenetic modifications are often reversible, and thus it is promising to improve ICT responses via epigenetic interference. A recent study showed that germline variants in prostate cancer patients have an impact on tumor epigenetic profiles: many germline variants influence methylation levels in prostate tumors but not in normal tissues [73], in a population-wide manner (Fig. 1). These results suggest that germline variants modulate DNA methylation and tumor epigenome to contribute to tumorigenesis and metastasis. Prostate cancer is a highly inheritable cancer [74], therefore, the links between germline variants and tumor epigenome in prostate cancer may not apply to other cancers.
GERMLINE PATHOGENIC VARIANTS MODULATE CANCER IMMUNE EVASION AND IMMUNOTHERAPY RESPONSE IN SUBSETS OF CANCER PATIENTS

Except for the population-wide influence of cancer immune-evasion and immunotherapy response by germline pathogenic variants, it has been found that some germline genetic factors have an impact on cancer immune-evasion and immunotherapy response in subsets of patients.

Germline variants in DNA-repair machinery genes regulate immunotherapy response

Dysfunction in DNA-repair machinery is another important contributing factor for cancer development, progression, and metastasis [75]. For a small fraction of cancer patients, their germline genomes contain pathogenic variants in the DNA repair genes, which lead to increased genome instability. Genome instability is a major driving force for increasing somatic mutations in tumor cells [39]. If a tumor has more somatic mutations, it might have more neoantigens being presented onto the cell surface and thus have a higher chance of responding to ICT. Several studies have shown that tumor mutational burden (TMB) is positively correlated with the response rate of ICT [76]. Currently, TMB has been used as a biomarker to predict ICT response [77], although the accuracy is not very high.

Germline variants in chromosomal instability genes contribute to cancer immune evasion

Chromosomal instability (CIN) is another hallmark for cancer [1]. Through an analysis of germline genomes of cancer patients from TCGA, we found that CIN is significantly enriched in the germline genomes of a small fraction of cancer patients (of 12 common cancer types) than that of non-cancer individuals (unpublished data). These results agree with the estimation that 60%–80% of human tumors exhibit chromosomal abnormalities [78,79]. Chromosome segregation errors (i.e., for CIN) can activate innate immune signaling through the introduction of genomic double-stranded DNA (dsDNA) into the cytosol and engagement of the cGAS-STING cytosolic dsDNA-sensing, anti-viral pathway [80,81]. This might explain CIN’s role in immune evasion, tumor evolution, and metastasis.

Germline variants in immune-related genes affect ICT response

In current ICT, CTLA4 is one of the key immune checkpoints. Inherited germline pathogenic variants in CTLA4 gene display a different response to ICT compared to wild-type CTLA4 [82,83]. In another study, it was found that germline mutations in CTLA4 would decrease numbers as well as functions of T-regulatory (Treg) cells, and lead to hyperproliferation of T cell and a progressive decrease in B cells [84]. Germline pathogenic variants in genes related to interleukins and chemokines show a different response to both anti-PD-1 and anti-CTLA4 treatments [85]. IFN signaling pathway is critical for tumor cells escaping. Germline pathogenic variants in IFN regulatory factor-5 (IRF5) are associated with different efficacy to immunotherapy [86,87]. Fc-γ receptors (FcγRs) are found on the surface of many immune cells such as NK cells and B cells, and function in regulating the immune system. Germline pathogenic variants in the FcγR gene compromised the efficacy of anti-CTLA4 treatment [88]. Germline pathogenic variants that could activate JAK3 led to increased expression of PD-L1 on both tumor cells and macrophages [89]. These results implied that germline variants in immune-related genes affect ICT response.

Germline pathogenic variants in the genes of immune cells modulate immune-evasion and ICT response

The human immune system includes more than 20 types of immune cells which are key members of innate and adaptive immune components and signaling molecules such as chemokines and cytokines. Germline pathogenic variants of TLR4, P2RX7, and FPR1 may modulate anti-tumor immune response [90,91]. Germline variants in genes encoding chemokines and their receptors, cytokines or molecules in IFN pathway, such as IL-10, CCL5, JAK, and IFNGR, could modulate the recruitment of immune cells into TIMEs [92–95]. Germline variants in HLA-E, which is recognized by specific NK cell receptors, affect ICT response [96]. Also, germline variants in genes in immune cells such as T cells, B cells, NK cells or macrophages could modulate immune-evasion and ICT response [64] (unpublished results).

IMPLICATIONS AND FUTURE DIRECTIONS

It is both interesting and unexpected that recent studies highlight an important role of germline genomes in regulating cancer immunity and immunotherapy response. The germline genome can restrict cancer somatic mutations through either HLA genotypes or functional/evolutionary constraints in a population-wide manner. In addition, germline genomes could encode genetic factors that shape tumor epigenomic events, oncogenic pathways, tumor recurrence, and cancer risk by
modulating cancer immunity. Moreover, germline variants regulate functions of immune cells such as NK cells to modulate cancer immunity and ICT response. The uncovering of the causal relationships between germline genomes, epigenome, somatic mutations in tumors and NK cell variability for regulating TILs in TIMEs would provide new avenues such as reprogramming TIMEs for cancer treatment.

Resolving of a long-term debate in cancer biology

It has been debated for many years whether genetic, extrinsic factors [97] (i.e., exposure to pathogen infection and environmental agents contributes to tumorigenesis) or intrinsic factors [98] (“bad luck hypothesis” — random error mutations during DNA replication in some stem cells could explain the majority of cancer population) contribute to tumorigenesis and metastasis. Although both “bad luck” and “extrinsic factor” hypotheses disagree with each other in terms of intrinsic and extrinsic factors for contributing tumorigenesis, they agree that heredity plays a minimal role in tumorigenesis. Indeed, from the cancer cell perspective, family history and a handful of known germline pathogenic variants may only explain less than 5% of cancer cases. Based on systems biology, in the cancer hallmark network framework, we proposed that pre-existing genome (i.e., in-born genome or germline genome) could play a dominant-heritable role in shaping tumorigenesis and metastasis [39]. The key limitation of both “bad luck” and “extrinsic factor” hypotheses is that they are cancer cell-centered and ignored the variance of the host immune system. Here we and others have shown that germline pathogenic variants play a significant role in the cancer-immunity system (Fig. 2), highlighting that heredity does not play a minimal, but dominant role in tumorigenesis. In summary, from the view of cancer cells, heredity plays a minimal role in tumorigenesis, however, from the view of the cancer-immunity system, heredity plays a critical role in both tumorigenesis and metastasis.

The necessity to integrate germline genomic information into the personalized immunotherapy

In the past, germline pathogenic variants have been ignored in both cancer research and clinics. With the new findings of germline genetic contributions to cancer-immunity, the factors influencing ICT response should be considered in an integrated manner — both germline and somatic mutations (including other genomic and epigenetic alterations such as DNA methylation) should be associated with clinical outcomes. From the evidence above, germline genetic background might affect the degree of tumor immune heterogeneity (TILs and TIMEs in tumors) and then affect ICT response. We have shown that germline pathogenic variants are able to predict prognosis, thus, it would be critical to integrate germline data with tumor data such as gene expression, genomic

Figure 2. Contributions of germline genomes (i.e., heredity), bad luck (i.e., replicate) and external factors (i.e., environmental) to tumorigenesis and metastasis. The degree of contributions for each factor to tumorigenesis and metastasis in each organ (bone, bone marrows, brain, thymus, respiratory tract, lung, stomach, colon, kidney, liver, ovary, and urinary system) is illustrated by the color depth (from light pink to deep red).
and epigenetic alterations to construct machine learning models to select patients for immunotherapy.

Both research and clinical applications of immunity in cancer are still in the early phase. Lots of questions remain unanswered. For example, how many cancer immune evasion strategies can tumors use? Does a tumor utilize only one or several immune-evasion strategies? Which are the key immune evasion mechanisms that might be modulated by germline pathogenic variants? There are more questions which need to be asked and answered.

COMPLIANCE WITH ETHICS GUIDELINES

The authors Xue Jiang, Mohammad Asad, Lin Li, Zhanpeng Sun, Jean-Sébastien Milanese, Bo Liao and Edwin Wang declare that they have no conflict of interests.

This article is a review article and does not contain any studies with human or animal subjects performed by any of the authors.

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