Personalization of medical treatments in oncology: time for rethinking the disease concept to improve individual outcomes

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Abstract
The agenda of pharmacology discovery in the field of personalized oncology was dictated by the search of molecular targets assumed to deterministically drive tumor development. In this perspective, genes play a fundamental “causal” role while cells simply act as causal proxies, i.e., an intermediate between the molecular input and the organismal output. However, the ceaseless genomic change occurring across time within the same primary and metastatic tumor has broken the hope of a personalized treatment based only upon genomic fingerprint. Indeed, current models are unable in capturing the unfathomable complexity behind the outbreak of a disease, as they discard the contribution of non-genetic factors, environment constraints, and the interplay among different tiers of organization. Herein, we posit that a comprehensive personalized model should view at the disease as a “historical” process, in which different spatially and timely distributed factors interact with each other across multiple levels of organization, which collectively interact with a dynamic gene-expression pattern. Given that a disease is a dynamic, non-linear process — and not a static-stable condition — treatments should be tailored according to the “timing-frame” of each condition. This approach can help in detecting those critical transitions through which the system can access different attractors leading ultimately to diverse outcomes — from a pre-disease state to an overt illness or, alternatively, to recovery. Identification of such tipping points can substantiate the predictive and the preventive ambition of the Predictive, Preventive and Personalized Medicine (PPPM/3PM). However, an unusual effort is required to conjugate multi-omics approaches, data collection, and network analysis reconstruction (eventually involving innovative Artificial Intelligent tools) to recognize the critical phases and the relevant targets, which could help in patient stratification and therapy personalization.

Keywords Predictive preventive personalized medicine (PPPM) · Systems biology · Polypharmacology · Critical transitions

Introduction
Personalized medicine (PM) [1] — “to match the right drugs to the right patients” — has become a widely used term in both the scientific community as well as in the public debate, its vagueness notwithstanding. Indeed, PM lacks an unambiguous definition and is open to interpretation [2]. In the early
Advances in genomics have allowed to stratify patients into distinct groups, based on a few molecular differences, which are deemed to play a critical role in the pathogenesis process. According to these premises, disease individuation would principally rely on the gene-expression pattern associated in a specific patient, allowing depicting a one-to-one correspondence upon a hypothetical Cartesian space, between the “genomic signature” and the illness in each individual [7].

This Promethean dream comes true in 1999, when Francis Collins established in a seminal paper the ways the human genome would be used to predict, prevent, and treat disease in the next 10 years, so as a complete transformation in medical practice would be expected even before 2020 [8]. This is the (bewildering) promise. However, we are in 2020 and this astonishing revolution in medicine has not yet been glimpsed.

Clinical randomized trials have provided little if any evidence of benefits when patients with different diseases have been treated with personalized-based treatments [9, 10]. This failure has contributed to the emergence of the so-called “reproducibility crisis” in biology and medicine [11, 12]. Still worse, these disappointing results prompted to cast on doubt the foundational assumptions of precision medicine [13].

Indeed, even “simple,” monogenic diseases — in which a single point mutation is recognized as the “main” causative factor — are not “simple” in their pathogenesis, as a number of additional factors, distributed across different hierarchical levels of the living organisms (from the DNA to physiological apparatus), are ultimately responsible of the disease phenotype. A paradigmatic case is provided by sickle cell anemia, a classic monogenic disorder, in which the interplay among a number of context-dependent cues enacts the emergence of no less than six different pathological phenotypes [14]. Some patients develop principally painful crises with or without bony infarcts; others are prone to hemolytic emergencies; some develop vaso-occlusive crises, including stroke; still others develop acute chest syndrome, while many are phenotype normally, except for mild anemia. What matters here is that the treatment — to be “precise” and “personalized” — must be tailored according to the emerging clinical pathophenotype, and not based on the primary point mutation. In this case, PM could hardly fit the specific needs of each patient, given that the genotype does not match with the (disease’s) phenotype [15]. Additionally, gene variants (including mutated genes), initially thought to play a “pathogenetic” role in several complex diseases, have been “reclassified,” given that their involvement becomes “problematic” [16]. Overall, in many pathological conditions, gene variance does not seem to play a relevant role, as the genes involved do not seem either to possess any biological link with the pathogenetic mechanism, or they can offer any clinical utility for prognosis [17]. As a fact, the relative risks for the vast majority of gene variants rarely exceed 1.5, and these variants have added little useful predictive power to traditional risk prediction algorithms. Even for diagnostic purposes, wide genome analysis often fails to equate the predictive power of classical medical parameters (anamnesis, neighborhood, socioeconomic status, dietary habits), and sophisticated assessment of gene-expression patterns adds little
(if any) to conventional predictive models [18]. Consequently, the main obstacle to progress in PPPM approaches for cancer is the lack of validated prognostic and predictive biomarkers [19].

It has been argued that the discouraging results obtained by PM-based clinical trials should be attributed to selection bias and uncertainty in defining clear outcomes [20], while developing more sophisticated approaches to identify specific patients subsets — using broad molecular testing and integrated genomic data from liquid biopsy samples [21, 22] — would in principle help in overcoming such a failure. This is wrong, as although the number of patients eligible for genome-driven treatment has increased over time, these “tailored” drugs have helped only a minority of patients with advanced cancer [23]. Moreover, unambiguously statistical criteria for patient’s selection and outcome parametrization are still inadequate. Several technical methodologies — including unsupervised discovery and data mining — have been used without explicating clear hypotheses to justify the observed (statistical) correlations. Therefore, selection bias and factors that can distort exposure-outcome correlation are usually overlooked. However, population-based studies of a disease require specific theoretical assumptions that inform data collection and allow to ascertain both exposures and outcomes in a standardized fashion [24].

To address such issues, instead of reconsidering the biological assumptions on which the PM strategy has been developed, an increasing number of scientists preferred to bypass that hurdle by adopting a new statistical-biometric approach as such provided by the Big Data Theory. Namely, the development of omics and systems biology has promoted one to gradually change paradigms in oncology from traditional single-factor strategy to multi-parameter systematic strategy [25].

Yet even this framework showed to be unsuccessful [26]. Data handling does not produce any new information by itself, as correlation does not mean “causation.” Furthermore, few prognostic factors or systems are robustly validated, and still fewer have made a convincing difference in health outcomes or in prolonging life expectancy [27]. In most diseases and outcomes, a considerable component of the prognostic variance remains unknown for our understanding of the critical mechanisms on whom the disease process depends is still insufficient. Additionally, most correlations are spurious, i.e., very large databases likely contain arbitrary correlations [28]. Empowerment of statistical analysis and sophisticated modeling cannot compensate for the lack of theory into which information from experiments need to fit. Computationally intensive tools for the exploitation of huge data sets are still based on poorly designed model; presumptively, they can only help in generating new hypotheses, but not true explanations. Consequently, applications of Big Data Theory have met with limited success in scientific domains, up to now [29]. Thereby, a new theoretical framework is urgently warranted “as a guide to experimental design for maximal efficiency of data collection and to produce reliable predictive models and conceptual knowledge” [30].

Inadequacies of theoretical assumptions at the root of the personalized medicine

Especially in the oncologist community, several scholars have enthusiastically welcomed personalized medicine (i.e., precision medicine) as a solution to find selective drugs able to modulating sensitive, key targets deemed to “drive” the overall process of tumor regression [31, 32]. Target-based therapies are broadly rooted into a theoretical framework such as provided by the Somatic Mutation Theory (SMT).

According to SMT, cancer is a cell-based disease [33], due to the accumulation of somatic mutations and/or chromosomal aberrations that alter the control of proliferation in a single cell that eventually will generate a neoplasia. This approach is essentially “reductionist” in essence, as it posits that the system — cells, tissues, cancer — can be explained by studying its parts in isolation, while the principal causative factor must be identified at the lowest level of organization (i.e., DNA, proteins, and so forth). This model has been extensively criticized given that is unable to accommodate with an increasing number of controversial and paradoxical results [34, 35]. For instance, accumulating evidence shows that genomic alterations, such as those in BRAF, RAS, EGFR, HER2, FGFR3, PIK3CA, TP53, CDKN2A, and NF1/2 genes — all of which are considered hallmark drivers of specific cancers — can also be identified in benign and premalignant conditions, occasionally at frequencies higher than in their malignant counterparts [36, 37].

Moreover, we usually forget that the search for critical genomic targets is seriously flawed from the outset by the unavoidable, intrinsic genomic heterogeneity of cancerous tissues [38]. Distinct mutations can be present as high as 100 million even in a single tumor [39], while sequencing and genome analysis of multiple biopsies from different regions of the same tumor reveals the wide spreading of genomic heterogeneity [40]. Intratumor heterogeneity is present since the early steps of cancer development and it is uncanny that chemotherapy can select further subclones, sharing increased aggressiveness and dedifferentiation, leading cancer to progressively becoming insensitive to any medical control [41].

Such heterogeneity yields a mosaic of different cells and, ultimately, can hamper cancer treatment [42]. This inescapable complexity contradicts the superficial textbook concept of clonal expansion of a dominant cell clone carrying the oncogenic mutation that “takes over” the entire tumor. In the real world, each cell shows a distinct set of mutations, while...
the subset of cells carrying “driver” mutations is unable to canalize the overall population into a unique, homogenous gene-expression pattern. In fact, although some tumors harbor a dominant population emerging from clonal selection, genomic diversity is the rule than the exception [43]. This process not only entails the topological distribution of cancer cells within the tumor, but also emerges at multiple time points during tumor progression, as demonstrated by liquid biopsies obtained from serial samples [44]. These data indicate that distinct clusters from the same tumor may undergo independent progression pathways, which culminate into the simultaneous presence of different phenotypic populations, each one harboring different malignant traits. This bewildering variety in cancerous phenotypes cannot be merely ascribed to differences in genomes but call into question the existence of epigenetic mechanisms and subtle modulation of the gene regulatory networks [45]. This non-genetic heterogeneity of tumor cell states defies a “precise” genotype–phenotype causal relationship and allows them to adapt to both environmental perturbations (nutrients availability, cells crowding, and hypoxia) as well as treatments without enacting the appearance of additional mutations. Again, these findings confirm that there is no straightforward linear causal relationship between tumor genotype and phenotype [46]. Moreover, cancer cells within individual tumors often exist in distinct phenotypic states. Given certain conditions, any subpopulation of cells (i.e., with different phenotypes) will return to equilibrium phenotypic proportions over time, after experiencing a critical transition [47], which enacted the disclosure of multiple, branching differentiating trajectories [48]. Notice that cancer stem-like cells arise de novo from non-stem-like cells, thus “regenerating” the malignant potentiality of the tumor. This is why seemingly identical cells respond differently to treatments, given that phenotypic and genotypic differences provide differentiated response by activating even opposite outcomes in cell behavior and ultimately escaping the drug-induced inhibition on specific pathways [49]. Overall, these findings highlight how complex and unstable is the gene-expression pattern of a tumor population, within each patient [50].

This body of evidence cannot easily accommodate with the prevailing carcinogenesis model. Instead, accumulated pitfalls contribute to laying bare the inadequacy of the SMT. To overcome these limitations, SMT became subject to a number of course-corrections, which strive to integrate new concepts by recurring to twisted arguments [51]. Latest versions of SMT include even the microenvironment — viewed as instrumental in promoting carcinogenesis — while trying to preserve the native mutation-based hypothesis [52]. All in all, these attempts look like the epicycle-based strategy used in ancient time to accommodate with the experimental facts that challenged the Ptolemaic system [53]. In alternative, in order to get rid of these conundrums, cancer has been proposed as an emergent phenomenon, due to a deregulated cross talk between cells and their microenvironment [54]. Carlos Sonnenschein and Ana Soto have conceptualized this new framework within the Tissue Organization Field Theory (TOFT) [55, 56]. TOFT is anchored at the tissue level of biological organization and conceives the development of cancer as a relational problem, focusing not on a single cell type but, as in organogenesis, on the interactions among different cell types and their microenvironments. The alteration in the interplay among those components involves different levels of organization and a number of secondary changes, eventually including the emergence of disrupted gene-expression patterns. In agreement with TOFT, mutated genes are the result, and not the cause, of the disrupted normal tissue architecture that eventually ends up in fostering cancer onset [57]. Therefore, changes in genomic profiles or in biochemical pathways can only be “associated” rather than considered as “causative.” Admitting this bitter conclusion would deprive precision medicine in oncology of its rationale, given that PM relies on targeting a validated and genetically stable driver of disease. To date, proof-of-concept trials have not supported this premise [12, 58]. Moreover, target-based treatments in oncology suffer from two major drawbacks: (1) Currently available inhibitors of specific pathways provide only minimal or complete blockade of biochemical pathways and are therefore inefficient or too toxic to be used [59]. (2) Second, critical networks in cancer — as well as in living cells — show a bewildering plasticity and adaptability, even under harsh environmental conditions, thus allowing the system to escape from programmed cell death [60]. Overall, those considerations help in explaining the shift in interest from the cancer cell to the stroma [61] and substantiate the relevance of microenvironment-based studies in search of new treatment options, as advocated by TOFT [62].

Personalized treatments are unable to cope with such an overwhelming complexity [63], and the small improvement in cancer survival recorded in the last years can only minimally be ascribed to target-based therapy [64, 65]. Namely, randomized, large studies with different combinations of target-based treatments in a number of cancer types failed to demonstrate any significant encouraging efficacy in any of the treatment arms or patient subsets [66, 67]. For instance, a multicenter randomized trial of treatment based on tumor sequencing compared with conventional cancer treatment showed no advantage of sequencing [68], as did the NCI-MATCH (National Cancer Institute–Molecular Analysis), in which almost 6,000 patients have been enrolled [69]. Similarly, a basket trial testing molecularly guided treatment approaches for multiple mutations in advanced non–small-cell lung cancer was demonstrated to be unsuccessful [70]. Expectancies from the recently introduced immunotherapy approaches have been disappointing as well. Immunotherapy
had only limited effects on the drop in overall cancer mortality. The undisputed benefits for melanoma and metastatic lung cancer are impressive, but so far they affect relatively few people and are associated with life-threatening side effects [71]. Overall, the weight of this evidence prompted to suggest that “our best weapons against cancer are not magic bullets” [72]. Therefore, precision medicine — especially in oncology — has lost most its fashion, while raising embarrassing concerns. A sober view of the evidence derived from prospectively designed trials of personalized medicine inevitably leads us to consider “that our current oncology community will be guilty of hubris and of overpromising what we can deliver in a realistic time line” [73].

**Disease: how to reframe the concept**

It is disheartening that the debate on PM has left aside a preliminary premise: what do we mean by “disease”? [74] No doubt that a number of inconsistencies of our modern treatment strategies — including those claimed by PM — should be attributed to the controversial concept of human disease. The debate on that subject mostly developed around two opposite positions represented by Constructivism and Naturalism. The former essentially denies the naturalist thesis that disease necessarily encompasses bodily malfunction and claims for a general “reformation” of the concept of disease, conceived as a “societal,” historical-based construct [75]. On the contrary, according to the naturalistic perspective — by far the only to which the daily medical practice actually relies — a disease involves malfunction of organs/apparatus, which either can or cannot be perceived as such, i.e., by complaining symptoms [76].

The latter model originates from Virchow’s conclusion that all diseases result from cellular abnormalities [77]. Since the discovery of the double helix in the 50 s, however, that framework was strongly superseded by an even more reductionist approach, as that provided by the New Genetics, which posits that every disease can be traced back to the malfunctioning of a discrete number of genes [78]. Briefly, this reductionist model relies on the following three premises: (1) the disease recognizes a dominant (molecular) cause; (2) medical signs and symptoms — which altogether constitute the disease phenotype (the “pathophenotype”) — are linearly correlated with the molecular cause; (3) removal/correction of the underlying, putative “cause” will restore healthy conditions. Sad to say, that model still awaits to be vindicated beyond any reasonable doubt, especially for degenerative diseases or mental illness [79]. Moreover, such framework becomes problematic when considering in the perspective of “preventive” medicine [80]. Are presumptive markers of a “future” disease condition reliable enough to ask for a “preventive cure”? Could a genomic profile allow drawing a reliable probabilistic ascertainment of a future disease? And then, could someone with a “genetic predisposition” (whatsoever this really means) be considered already sick?

Nevertheless, in the last 30 years, diseases have been increasingly “equated” to the malfunctioning of a few, critical pathways or of their related driver genes. Consequently, drug discovery has been dominated by reductionism, aiming to identify drugs that activate or inhibit specific molecular targets. Unfortunately, therapeutic approaches based on such a simplistic paradigm often showed either unforeseen toxicity or lack of efficacy when tested in clinical trials [81].

Consequently, the adoption of this reductionist-based approach progressively distorted and shaped medical practice — namely by radically modifying the diagnostic methodology and the doctor–patient relationship — leading toward a new “disease taxonomy” [82]. Consequently, we are witnessing a number of inadequacies in the current medical practice, which often reflect a lack of specificity (i.e., inability in defining a disease unequivocally), and a lack of sensitivity (i.e., incapacity in recognizing preclinical, true causative state of disease). Ultimately, this model is proven to be confounding, as it often posits wrong correlations between the disease-associated biological parameters (usually identified only when illness reaches a “stable-state”) and the alleged causative processes, thereby prejudicing efficient treatment strategies.

At a first glance, the starting premise of PM relies on the following, abridged statement: one genotype-one pathophenotype. This is wrong, even for simple monogenic diseases. The abovementioned example of sickle cell anemia highlighted a complex interplay of different causative factors — spanning from cell to higher levels of living system’s organization — which ultimately fosters the emergence of no less than six different disease phenotypes. Similarly, EBV infection is recognized to promote mononucleosis or Burkitt’s lymphoma.

In both situations, B cells represent the primary target of EBV [83]. In Western countries and in presence of well-regulated immune system, the virus-induced proliferation of B cells is intrinsically controlled by CD8+ and CD4+ T cells. In this setting, the EBV infection generally triggers infectious mononucleosis. On the contrary, in inhabitants of the several equatorial regions of the world, EBV induces the development of lymphoid tumors [84]. The difference lies on the previous medical history of the host and on the specific interactions that EBV triggers with the immune system of each individual [85]: a set of factors that cannot be deduced from the genomic analysis neither of the viral genome, nor the genome of the host. In simple words, the specific disease arises from a dynamic process in which the primary causative factor (the virus) interacts in a non-linear fashion with complex apparatus of the host.
(not limited to the cellular compartment); organs, cells, and tissues are in turn “shaped” and “customized” by the previous medical history of the organism living in a very unique environment [86]. As a result, EBV infection is currently known as the main “causative” factor of a number of disparate diseases, including pharyngeal carcinomas [87], gastric cancer [88], and non-malignant illness, such as the childhood disorders of Alice in Wonderland Syndrome [89], systemic lupus erythematosus [90], and acute cerebellar ataxia [91]. This example epitomizes that a disease cannot be considered a “static” state but should instead be considered a dynamic process, ruled by non-linear dynamical relationships, which can ultimately drive the system toward very different outcomes. Broadly speaking, “genetic defects” cannot predict the pathophenotype, which ultimately emerges from the complex interactions among different factors, distributed across several, hierarchically organized levels.

According to the PM premises, the causative factor(s) that are thought to contribute to the disease process still should be “at work” at the time of treatment. However, as happen for several conditions, the mechanism/gene responsible for the onset of the illness might have exerted its action during early pathogenic steps and could no longer be active during the steady state of the disease, when diagnosis is usually reached. Some developmental-based diseases, like mental illness [92] or cardiovascular diseases, fall within this category, as well as some cancers that “must” lose their mutated “driver” oncogenes precisely when they metastasize and — paradoxically — become more aggressive [93].

However, pathogenic interactions are distributed across a space-temporal continuum, given that a number of genomic-related factors are likely to act only during some critical developmental phases (during the intra-uterine life, at the birth, in the neonatal, and pre-pubertal period). Therefore, disease development is a time-dependent process, tightly linked to the patient’s history. In the last resort, disease should be viewed as a manifestation of developmental plasticity, the phenomenon by which one genotype can give rise to a range of different physiological or morphological states in response to different environmental conditions during development [94]. Therefore, numerous chronic diseases are currently supposed to arise from some “disturbances” acquired during critical developmental periods [95].

Epigenetic and post-translational changes can efficiently begin as early as during pregnancy (still in the womb) and may be affected by the paternal and maternal environments, as well as by early life events (dietary habits, childhood diseases, premature exposure to environmental carcinogens and toxicants like endocrine disruptors) [96]. These modifications play a critical role in shaping cells and tissue sensitivity to carcinogens, ultimately favoring the emergence of cancer in the adult life [97, 98].

This premise carries additional consequences. Diseases — all together with their “causative” targets — are usually recognized by late-appearing manifestations. However, disease development entails several time-distributed steps, and specific treatments should be put in place at each distinct phase in which a specific rewiring of the Gene Regulatory Network is likely to occur (Fig. 1). Moreover, diagnostic

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**Fig. 1** Disease as a historical, dynamical process. Most of harmful stimuli (exogenous toxicants, microbes, metabolic factors, radiation, etc.) hit different kind of cells and tissues, as only very few pathogenic cues interact with a single cell type. The response can be appreciated at both local and organismal level, involving the participation of many different tissues and structures. Overall, this entrenched cooperativity contributes to reshaping the Gene Regulatory Network as well as several biochemical pathways. The entire process proceeds across different bifurcation points (A, B) displacing itself through different attractors (i.e., phenotypic states), before reaching a stable “disease-state”
parameters and putative causative factors are frequently associated with the steady state of the disease. This approach involves the obvious risk to consider a late-emerging symptom/target as the driver-causative element of the pathogenic process, while discarding early, critical signs.

Definitely, current PM-based treatments are unable to cope with such an overwhelming complexity, and their acknowledged failure in curing cancer cannot be viewed as an unannounced surprise [61] [63]. Consequently, “we overdiagnose, overtreat, and overpromise, with high costs and without clear benefits” [62].

Reconsidering the concept of human disease according to a systems biology approach

The reductionist model of disease overlooks the relevance of multifactorial etiology of the disease and underestimates the robustness and resilience of the (pathologic) phenotype, especially when pharmacologically perturbed. For instance, single-gene knockout or complete silencing has shown little contradictory or even null effect on the pathophenotype [99]. Inhibiting a selected pathway can be insufficient in controlling the “corresponding” biological function, as the network can switch toward alternative in response to changing requirements of the context in which the system belongs [100, 101]. The switch occurs at specific “tipping points” where the system can enter into previously “unknown” attractors, thus acquiring “unexpected” features, including the resilience to a wide range of perturbing factors [102]. It should be stressed that at the bifurcation point, we usually observe an increase in the fluctuation of several parameters. Fluctuations are critical for enzymes to work, for a receptor to switch between states, and for the chromatin to express the right protein at the right time [103]. Overlooking these fluctuations will likely affect the identification of those states in which the system affords the choice in between different cell fate commitment [103]. These considerations may have huge consequences, given that the identification of such parameters would allow in recognizing those targets that are instrumental in driving transitions in the pre-disease state, thus performing true “preventive diagnostic.” Anticipating the transition from the pre-disease to the stable disease state represents a testable “personalization” of medical treatment. In other words, target recognition in the preventive perspective should shift from the disease to the highly dynamic and complex pre-disease state.

The contribution of internal or environmental constraints in “driving” such transitions is mandatory, as they represent additional “causative factors” [104]. The emergence of a specific network associated with the time-dependent state of the disease process can be properly ascertained only if the specific microenvironmental field is concurrently contemplated. Genetic regulatory networks and the context-dependent constraints are tightly intertwined and therefore a successful therapeutic strategy should embrace all of them if the aim is properly to cure the patients, and not only “to fix” a “singled out” pathway [105]. Indeed, constraints exert a mandatory function in driving the systems toward distinct attractors, i.e., specific phenotypes recognizable by a different architecture of its Gene Regulatory Network (Fig. 2).

A comprehensive approach to this problem would enable in providing a complex network structure, constituted by modular sub-systems, whose (non-linear) interaction will drive the organism response toward emergent properties, i.e., disease or health. Therefore, human disease needs to be conceptualized as an “emergent property” of the human body [106]). Overall, these considerations ask for revisiting the concept of illness on which any personalized medicine should deal with. Systems biology may help in establishing a new model, able to integrate different “causative” factors, distributed across different scales (from molecules to organs) and times (from the early life to the present). Within this framework, response parameters should be provided by the overall system estimate, rather than on singled-out molecular target [107]. In other words, disease should be conceptualized as a non-linear dynamic process displaying classical features of complex systems, including resilience, sensitivity to initial conditions, and multi-attractor accessibility. Transition from different states — healthy, high-risk conditions, pre-disease, and disease states — has been documented and modeled in some instances [108, 109]. Namely, the existence of “critical period” during the lifespan (especially during early life) has been highlighted by studies that have identified a strong link between those periods and the appearance of different diseases in the adult period [110, 111]. During the development, organs and systems of the body go through “critical” periods, in which their sensitivity to internal and environmental perturbations is dramatically amplified [112]. While it is intuitive that such a plasticity — i.e., the capacity to dynamically respond to surrounding stresses by shaping morphology and functions — is an advantage from the evolutionary perspective, nonetheless such a capability exposes the organism to unwarranted risks. Indeed, when the system experiences important perturbations, even pathological states can arise and eventually be maintained if they become reliable, adaptive issues.

Briefly, the physiological system can travel across that metaphorical landscape (remnant of the Waddington’s approach) [113], accessing different attractors, depicted as different physiological/pathological states — a normal state, a pre-disease state, a disease state — in which the disease can alternatively move toward progression or healing. Identification of these bifurcation points could help in understanding the meaning of the overall process and in managing it toward beneficial outcomes [114]. Finally, identification of
biomarkers (“early-warning signals”) indicating an imminent bifurcation or sudden deterioration before the critical transition occurs can help in planning an appropriate management of the disease or the pre-disease state, thus providing the “preventive” strategy with an entirely new meaning [115].

Conclusions and expert recommendations

The practice of medicine is currently undergoing a paradigm shift, aimed at treating a disease by identifying the specific fingerprint displayed by the illness in an individual; insofar each patient can be featured through a personalized data-driven approach, as that allowed by the convergence of big data and multi-omics approaches.

Such a task entails several ethical, social, and regulatory issues [116, 117]. As a consequence, several theoretical and methodological premises on which PPPM relies require to be reconsidered given that the complexity of a bewildering number of extragenomic factors undermines the predictability of PM models based prevalently on genomic data [118, 119].

For most common diseases, hundreds of genetic risk variants with small effects have been identified, and it is hard to establish an unambiguous picture of who is really at “risk” and for “what.” Definitely, no convincing evidence allows in recognizing a one-to-one correspondence between genomic alterations and widespread, complex diseases for which “traditional” markers are much better predictors than extensive DNA data sets [118]. Notice that inappropriate and exaggerated expectations in PPPM promises will easily turn out into false hope. People believing not to be at risk after being reassured having performed a “genome-based test” will probably dismiss appropriate behaviors and health care precautions; on the contrary, people “informed” to be at high risk shall probably endorse a resigned attitude, therefore discarding medical controls and treatments [120].

Despite PPPM missed its promises, we yet continue to overinvest our hope (and money) in genomic-based approaches. Yet genetics cannot deliver unaffordable expectations — as recognized even by PM advocates during the recent Covid-19 pandemics [121] — given that genes do not constitute that “privileged” level of causation the reductionist medicine was searching for establishing a deterministic biological model [122]. In simple words, we actually lack the minimal facts needed to support the hope that genetics can be a key to realizing that vision. This statement implies we should orchestrate a stratified and multilevel diagnostic and treatment approach [123].

A more prudent reappraisal of PPPM has evidenced the paradox behind this approach. In principle, PM-based
treatments are themselves more precise than standard chemotherapeutic protocols although the clinical evidence supporting the benefits of these therapies is often considerably less precise and still awaits sound confirmation [124]. Paradoxically, clinical trials designed to assess PPPM efficiency demonstrated a lack of precision in respect to conventional trials [125]. Namely, these studies comprise several treatment arms in which far fewer patients than in conventional phase III investigations are generally enrolled. It is quite disturbing that these patients are usually not randomized owing to the difficulties in determining a sufficient number of eligible patients or in planning a proper control treatment. Furthermore, results are biased by the limited choice of endpoints (usually confined to the overall response rate), thus undermining the trials' validity.

Secondly, to “extract” useful knowledge from a complex system, one must focus on the right level of description [126]. No doubt that the disease is an emergent phenomenon involving the organism and not limited to molecules or cells. Thereby, any attempt to establish a precision-based treatment should look at the “system,” instead of focusing on single molecules or pathways. To sum up, despite undeniable performances recently performed in the context of predictive approaches, targeted prevention, and personalization of medical care (PPPM/3PM), several aspects are waiting for innovative solutions [127].

In detail, treatments should be tailored according to the “timing-frame” of each condition. This approach can help in detecting those critical transitions through which the system can access different attractors leading ultimately to diverse outcomes — from a pre-disease state to an overt illness or, alternatively, to recovery. Identification of such tipping points can substantiate the predictive and the preventive ambition of the PPPM/3PM.

Namely, analyses performed on liquid biopsies by means of mass spectrometry-based technologies [128] have allowed the identification of a wide array of biomarkers and of their specific pattern of expression that both can provide valuable diagnostic information for the identification of overt disease, its preliminary stages, and ultimately the very initial (reversible) conditions. Thorough investigation carried out on liquid biopsy specimens can be instrumental in drawing precise phenotyping, i.e., a multi-omics fingerprinting — that, in turn, would allow the creation of tailored treatment algorithms, thus providing the most optimal clinical approach for personalized, predictive, and preventive medical service [129].

This perspective implies that we should shift from targets to processes and identify a multi-target array — entailing different mechanisms of action — as suggested by the network polypharmacology approach [130]. Modulation of processes implies we should be able to “redraw” the disease-related landscape, favoring the system displacement from pre-clinical state or true disease-states toward healing pathways. Disease could eventually be “reverted,” involving also a “reprogramming” of the gene-regulatory network, as proposed in some carcinogenic models [131–133]. Indeed, as cancer can be successfully “reverted” through the modification of the dynamical cross talk with its microenvironment, interactive cell-stroma networks must be recognized as a target for pharmacological intervention [60, 134, 135].

How should we investigate the evolution of such a complex system over time? We know that distributed non-linear network systems are hardly mathematically tractable when matched with simple feedback systems, usually described by means of control theory. Disease response is habitually simplified by describing changes in a single (or a few) parameter. Instead, we have to move from target-related parameters to system parameters, which could capture those modifications that could likely impact on the whole systems dynamics [136].

At the core of this new strategy, a relevant role is sustained by multi-omics data, which can likely help in moving from a genome-centered to a phenome-centered research practice. However, an unusual effort is required to conjugate different high-throughput analytical methods, data collection, and network analysis reconstruction (eventually involving innovative Artificial Intelligent tools) to recognize critical phases and relevant targets [137]. Noteworthy, this approach could help in allowing a patient stratification according to a very different perspective [138, 139].

A relevant case in point is constituted by metabolomics studies that — in principle — can satisfy those needs. Metabolites fluctuations usually amplify subtle modulation of the genome/proteome networks over time and metabolomics can describe changes in complex systems dynamics as well as its “adaptive” phenotype in presence of a wide
range of perturbations [140–142], thus providing a new basis for establishing a proper PPPM/3PM framework. Noticeably, a striking correlation can be established between metabolomics and proteomic changes, and both can play a driving role during cell/tissues phenotypic transitions. Moreover, proteomic changes can be viewed as the ultimate output of several processes — transcriptomic variations, splicing, post-translational modifications, translocation/re-distribution, spatial conformation, and pathway-network systems — that collectively might contribute to clarify the mechanisms of a disease and to recognize therapeutic targets [143].

Future interventions should aim to find the way to “modulate” the metabolomic fingerprint of a disease in a “precise” and efficient fashion, instead of searching for hypothetical “causative target” and “magic bullets.” Some pioneering approaches demonstrated that such an approach could successfully recognize subsets of patients within the same disease, characterized by different pathophenotypes and distinct activated pathways that can be successfully targeted with very different treatments [144]. This selection is performed through a systems biology approach and does not rely on single targets as it focuses on the overall behavior of the system under scrutiny.

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Declarations

Ethics approval and consent to participate NA

Consent for publication All authors have given their consent to the publication.

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References

1. Hamburg MA, Collins FS. The path to personalized medicine. N Engl J Med. 2010;363(4):301–4. https://doi.org/10.1056/NEJMp1006304.

2. Evers AW, Rovers MM, Kremer JA, Veltman JA, Schalken JA, Bloem BR, van Gool AJ. An integrated framework of personalized medicine: from individual genomes to participatory health care. Croat Med J. 2012;53(4):301–3. https://doi.org/10.3325/cmj.2012.53.301.

3. Leonti M, Casu L. Traditional medicines and globalization: current and future perspectives in ethnomedicine. Front Pharmacol. 2013;4:9. https://doi.org/10.3389/fphar.2013.00092.

4. Fojo T. Precision oncology: a strategy we were not ready to deploy. Semin Oncol. 2016;43(1):9–12.

5. Letia A. Functional precision cancer medicine—moving beyond pure genomics. Nat Med. 2017;23(9):1028–35. https://doi.org/10.1038/nm.4389.

6. Snyderman R, Meade C, Drake C. Value of personalized medicine. JAMA. 2016;315(6):613. https://doi.org/10.1001/jama.2016.17136.

7. Joyner MJ, Prendergast FG. Chasing Mendel: five questions for personalized medicine. J Physiol. 2014;592(11):2381–8. https://doi.org/10.1113/physiol.2014.272336.

8. Collins FS. Shattuck lecture—medical and societal consequences of the Human Genome Project. N Engl J Med. 1999;341(1):28–37. https://doi.org/10.1056/NEJM199907013410106.

9. Prasad V, Fojo T, Brada M. Precision oncology: origins, optimism, and potential. Lancet Oncol. 2016. https://doi.org/10.1016/S1470-2045(15)00620-8.

10. West HJ. No solid evidence, only hollow argument for universal tumor sequencing show me the data. JAMA Oncol. 2016;2:717–8.

11. Ioannidis JP. Why most published research findings are false. PLoS Med. 2005. https://doi.org/10.1371/journal.pmed.0020124.

12. Bizzarri M. Is Biology in an existential crisis? A diagnostic analysis and perhaps… an effective treatment (Editorial). Organisms Journal of Biological Sciences. 2019;2(2):1.

13. Joyner MJ, Paneth N. Promises, promises, and precision medicine. J Clin Invest. 2019;129(3):946–8. https://doi.org/10.1172/JCI126119.

14. Kato GJ, Gladwin MT, Steinberg MH. Deconstructing sickle cell disease: reappraisal of the role of hemolysis in the development of clinical subphenotypes. Blood Rev. 2007;21(1):37–47. https://doi.org/10.1016/j.blre.2006.07.001.

15. Loscalzo J, Kohane I, Barabasi AL. Human disease classification in the postgenomic era: a complex systems approach to human pathobiology. Mol Syst Biol. 2007. https://doi.org/10.1038/msb4100163.

16. SoRelle JA, Thedeson DM, Arnold S, Gwotay G, Park JY. Clinical utility of reinterpreting previously reported genomic epilepsy test results for pediatric patients. JAMA Pediatr. 2019. https://doi.org/10.1001/jamapediatrics.2018.2302.
17. McClellan J, King MC. Genetic heterogeneity in human disease. Cell. 2010;141(2):210–7. https://doi.org/10.1016/j.cell.2010.03.032.
18. Morris RW, Cooper JA, Shah T, Wong A, Drenos F, Engmann J, McLachlan S, Jeffers B, Dale C, Hardy R, Kuh D, Ben-Shlomo Y, Wannamethee SG, Whincup PH, Casas JP, Komivaki M, Kurni M, Talmud PJ, Price JF, Dudbridge F, Hingorani AD, Humphries SE, UCL EB Consortium. Marginal role for 53 common genetic variants in cardiovascular disease prediction. Heart. 2016;102(20):1640–7. https://doi.org/10.1136/heartjnl-2016-309298.
19. Duffy M. Personalised treatment for cancer: role of biomarkers. EPMA J. 2014;5(Suppl 1):23a.
20. Schweizerle M, Zhao M, Lee J, et al. Association of biomarker-based treatment strategies with response rates and progression-free survival in refractory malignant neoplasms: a meta-analysis. JAMA Oncol. 2016;2(11):1452–9.
21. Lebofsky R, Decraene C, Bernard V, Kamal M, Blin A, Leroy Q, Rizo Frio T, Pierron G, Callens C, Bieche I, Saliou A, Madic J, Rouleau E, Bizard FC, Lantz O, Stern MH, Le Tourneau C, Pierga JY. Circulating tumor DNA as a non-invasive substitute to metastasis biopsy for tumor genotyping and personalized medicine in a prospective trial across all tumor types. Mol Oncol. 2015;9(4):783–90. https://doi.org/10.1016/j.molonc.2014.12.003.
22. Gyawali B. Point: The imprecise pursuit of precision medicine: are biomarkers to blame? J Natl Compr Canc Netw. 2017;15(7):859–62. https://doi.org/10.6004/jnccn.2017.0126.
23. Marquart J, Chen EY, Prasad V. Estimation of the percentage of US patients with cancer who benefit from genome-driven oncology. JAMA Oncol. 2018;4(8):1093–8. https://doi.org/10.1001/jamaoncol.2018.1660.
24. Joyner MJ, Paneth N. Seven questions for personalized medicine. JAMA. 2015;314(10):999–1000. https://doi.org/10.1001/jama.2015.7725.
25. Zhan X, et al. Multi-parameter systematic strategy opinion that predicts, prevents, and personalized treats a cancer. EPMA J. 2014;5(Suppl 1):A25.
26. Prasad V. Perspective: the precision-oncology illusion. Nature. 2016;537(7619):S63. https://doi.org/10.1038/537S63a.
27. Ioannidis JPA. Limits to forecasting in personalized medicine: an overview. Int J Forecast. 2009;25:773–83. https://doi.org/10.1016/j.ijfor ecast.2009.05.003.
28. Calude CS, Longo G. The deluge of spurious correlations in big data. Found Sci. 2017;22(3):596–612. https://doi.org/10.1007/s10699-016-9489-4.
29. Karpatne A, et al. Theory-guided data science: a new paradigm for scientific discovery from data. IEEE Trans Knowl Data Eng. 2017;29(10):2318–31. https://doi.org/10.1109/TKDE.2017.2720168.
30. CoveyEN, Dougherty ER, Higfield RR. Big data need big theory too. Philos Trans A Math Phys Eng Sci. 2016. https://doi.org/10.1098/rsta.2016.0153.
31. Kaelin WG Jr. The concept of synthetic lethality in the context of anticancer therapy. Nat Rev Cancer. 2005;5(9):689–98. https://doi.org/10.1038/nrc1691.
32. Gambardella V, Tarazona N, Cejalvo JM, et al. Personalized medicine: recent progress in cancer therapy. Cancers (Basel). 2020;12(4):1009. https://doi.org/10.3390/cancers12041009.
33. Weinberg RA. One renegade cell: how cancer begins. New York: Basic Books; 2008.
34. Bizzarri M, Cucina A, Conti F, D’Anselmi F. Beyond the oncogene paradigm: understanding complexity in cancerogenesis. Acta Biotheor. 2008;56(3):173–96. https://doi.org/10.1007/s10441-008-9047-8.
35. Weinberg RA. Coming full circle—from endless complexity to simplicity and back again. Cell. 2014;157(1):267–71. https://doi.org/10.1016/j.cell.2014.03.004.
36. Martincocorena I, Roshan A, Gerstung M, Ellis P, Van Loo P, McLaren S, Wedge DC, Fullam A, Alexandrov LB, Tubio JM, Stebbings L, Menzies A, Widaa S, Stratton MR, Jones PH, Campbell PJ. Tumor evolution High burden and pervasive positive selection of somatic mutations in normal human skin. Science. 2015;348(6237):880–6. https://doi.org/10.1126/science.aaa6806.
37. Kato S, Lippman SM, Flaherty KT, Kurzrock R. The conundrum of genetic “Drivers” in benign conditions. J Natl Cancer Inst. 2016. https://doi.org/10.1093/jnci/djw036.
38. Kamb A, Wec S, Lengauer C. Why is cancer drug discovery so difficult? Nat Rev Drug Discov. 2007;6(2):115–20. https://doi.org/10.1038/nrd2155.
39. Ling S, Hu Z, Yang Z, Yang F, Li Y, Lin P, Chen K, Dong L, Cao L, Tao Y, Hao L, Chen Q, Gong Q, Wu D, Li W, Zhao W, Tian X, Hao C, Hungate EA, Catenacci DV, Hudson RR, Li WH, Lu X, Wu CI. Extremely high genetic diversity in a single tumor samples to prevalence of non-Darwinian cell evolution. Proc Natl Acad Sci U S A. 2015;112(47):E6496–505. https://doi.org/10.1073/pnas.1519556112.
40. Gerlinger M, Horswell S, Larkin J, Rowan AJ, Salim MP, Varela I, Fisher R, McGranahan N, Matthews N, Santos CR, Martinez P, Phillimore B, Begum S, Rabinowitz A, Spencer-Dene B, Gulati S, Bates PA, Stamp G, Pickering L, Gore M, Nicol DL, Hazell S, Futreal PA, Stewart A, Swanton C. Genomic architecture and evolution of clear cell renal cell carcinomas defined by multiregion sequencing. Nature Genet. 2014;46(3):225–33. https://doi.org/10.1038/ng.2891.
41. Bhang HE, Ruddy DA, Krishnamurthy Radhakrishna V, Caushi JX, Zhao R, Hims MM, Singh AP, Kao I, Rakiec D, Shaw P, Balak M, Raza A, Ackley E, Keen N, Schlabach MR, Palmer M, Leary RJ, Chiang DY, Sellers WR, Michor F, Cooke VG, Korn JM, Stoege ME. Studying clonal dynamics in response to cancer therapy using high-complexity barcoding. Nat Med. 2015;21(5):440–8. https://doi.org/10.1038/nm.3841.
42. Seoane J. Cancer: Division hierarchy leads to cell heterogeneity. Nature. 2017;549(7671):164–6. https://doi.org/10.1038/ nature23546.
43. Navin N, Krasnimtz A, Rodgers L, Cook K, Meth J, Kendall J, Riggins M, Eberling Y, Trogate J, Grubor V, Levy D, Lundin P, Márí S, Zetterberg A, Hicks J, Wigler M. Infiltrating tumor progression from genomic heterogeneity. Genome Res. 2010;20(1):68–80. https://doi.org/10.1101/gr.099622.109.
44. Ma M, Zhu H, Zhang C, Sun X, Gao X, Chen G. “Liquid biopsy”-ctDNA detection with great potential and challenges. Ann Transl Med. 2015. https://doi.org/10.1039/j.jissn.2305-5839.2015.09.29.
45. Brock A, Chang H, Huang S. Non-genetic heterogeneity—mutation-independent driving force for the somatic evolution of tumours. Nat Rev Genet. 2009;10(5):336–42. https://doi.org/10.1038/nrg2556.
46. Brock A, Huang S. Precision oncology: between vaguely right and precisely wrong. Cancer Res. 2017;77(23):6473–9. https://doi.org/10.1158/0008-5472.CAN-17-0448.
47. Gupta PB, Fillmore CM, Jiang G, Shapira SD, Tao K, Kuperwasser C, Lander ES. Stochastic state transitions give rise to phenotypic equilibria in populations of cancer cells. Cell. 2011;146(4):633–44. https://doi.org/10.1016/j.cell.2011.07.026.
48. Anderson K, Lutz C, van Delft FW, Bateman CM, Guo Y, Colman SM, Kemptser H, Moorman AV, Titley I, Swansbury J, Kearney L, Enver T, Greaves M. Genetic variegation of
clonal architecture and propagating cells in leukaemia. Nature. 2011;469(7330):356–61. https://doi.org/10.1038/nature09650.
49. Li RX, Zeng R. Dynamic proteomics for investigating the response of individual cancer cells under drug action. Expert Rev Proteomics. 2009;6(1):19–21. https://doi.org/10.1586/14789450.6.1.19.
50. Swanton C. Intratumor heterogeneity: evolution through space and time. Cancer Res. 2012;72(19):4875–82. https://doi.org/10.1158/0008-5472.CAN-12-2217.
51. Wallace DC. Mitochondria and cancer. Nat Rev Cancer. 2012;12(10):685–98. https://doi.org/10.1038/nrc3365.
52. Bissell MJ, Hines WC. Why don’t we get more cancer? A proposed role of the microenvironment in restraining cancer progression. Nat Med. 2011;17(3):320–9. https://doi.org/10.1038/nm.2328.
53. Sonnenschein C, Soto AM. And yet another epicycle. BioEssays. 2006;28(1):100–1. https://doi.org/10.1002/bies.20348.
54. Bussard KM, Boulanger CA, Booth BW, Bruno RD, Smith GH. Molecularly targeted therapy based on tumour molecular profiling versus conventional therapy for advanced cancer (SHIVA): a multicentre, open-label, proof-of-concept, randomised, controlled phase 2 trial. Lancet Oncol. 2015;16(13):1324–34. https://doi.org/10.1016/S1470-2045(15)00188-6.
55. DeLorenzo JV, Soto AM. Lethal and less lethal: diseasome perturbations lead to cancer. Epigenetics. 2016;11(3):244–58. https://doi.org/10.4161/epi.35313.
56. Bizzarri M, Cucina A. The society of cells: cancer and control of cell proliferation. New York: Springer Verlag. The Society of Cells – Cancer and control of cell proliferation. 1999.
57. Bizzarri M, Cucina A. SMT and TOFT: Why and how they are opposite and incompatible paradigms. Acta Biotheor. 2016;64(3):221–39. https://doi.org/10.1007/s10441-016-9281-4.
58. Sonnenschein C, Soto AM. Over a century of cancer research: inconvenient truths and promising leads. PLoS Biol. 2020. https://doi.org/10.1371/journal.pbio.3000670.
59. Bowen A, Casadevall A. Increasing disparities between resource inputs and outcomes, as measured by certain health deliverables, in biomedical research. Proc Natl Acad Sci U S A. 2015;112(36):11335–40. https://doi.org/10.1073/pnas.1504955112.
60. Greenwell IB, Ip A, Cohen JB. PI3K inhibitors: understanding toxicity mechanisms and management. Oncology (Williston Park). 2017;31(11):821–8.
61. Johnson GL, Stuhlmler TJ, Angus SP, Zawistowski JS, Graves LM. Molecular pathways: adaptive kinome reprogramming in response to targeted inhibition of the BRAF-MEK-ERK pathway in cancer. Clin Cancer Res. 2014;20(10):2516–22. https://doi.org/10.1158/1078-0432.CCR-13-1081.
62. Tchou J, Conejo-Garcia J. Targeting the tumor stroma as a novel treatment strategy for breast cancer: shifting from the neoplastic cell-centric to a stroma-centric paradigm. Adv Pharmacol. 2012;65:45–61. https://doi.org/10.1016/B978-0-12-397927-8.00003-8.
63. Bizzarri M, Cucina A. Tumor and the microenvironment: a chance to change the paradigm of carcinogenesis? Biomed Res Int. 2014. https://doi.org/10.1155/2014/934038.
64. Scannell JW, Bosley J. When quality beats quantity: decision theory, drug discovery, and the reproducibility crisis. PLoS ONE. 2016. https://doi.org/10.1371/journal.pone.0147215.
65. Hawkes N. High cost of cancer treatment doesn’t reflect benefits, say specialists. BMJ. 2011. https://doi.org/10.1136/bmj.d6220.
66. Bizzarri M. Do new anticancer drugs really work? A serious concern. Organisms. Journal of Biological Sciences. 2017(1):9–10.
67. Dingemans AM, Mellemra WW, Groen HJ, et al. A phase II study of sorafenib in patients with platinum-pretreated, advanced (Stage IIb or IV) non-small cell lung cancer with a KRAS mutation. Clin Cancer Res. 2013;19(3):743–51.
68. Papadimitrakopoulou V, Lee JJ, Wistuba II, Tsao AS, Fossella FV, Kalhor N, Gupta S, Byers LA, Izzo JG, GETTERING SN, Goldberg SB, Tang X, Miller VA, Skoulidis F, Gibbons DL, Shen L, Wei C, Diao L, Peng SA, Wang J, Tam AL, Coombes KR, Koo JS, Mauro DJ, Rubin EH, Heymach JV, Hong WK, Herbst RS. The BATTLE-2 study: a biomarker-integrated targeted therapy study in previously treated patients with advanced non-small-cell lung cancer. J Clin Oncol. 2016;34(30):3638–47. https://doi.org/10.1200/JCO.2015.66.0084.
69. Le Tourneau C, Delord JP, Gonçalves A, Gavoille C, Dubot C, Isambert N, Campone M, Trédan O, Massiani MA, Mauborgne C, Armanet S, Servant N, Bieche I, Bernard V, Gentien D, Jezequel P, Attignat V, Boyault S, Vincent-Salomon A, Servois V, Sabin MP, Kamal M, Paolletti X, SHIVA investigators. Molecularly targeted therapy based on tumour molecular profiling versus conventional therapy for advanced cancer (SHIVA): a multicentre, open-label, proof-of-concept, randomised, controlled phase 2 trial. Lancet Oncol. 2015;16(13):1324–34. https://doi.org/10.1016/S1470-2045(15)00188-6.
70. Eckhardt SG, Lieu C. Is precision medicine an oxymoron? JAMA Oncol. 2019;5(2):142–3. https://doi.org/10.1001/jamaoncol.2018.5099.
71. Tchou J, Conejo-Garcia J. Targeting the tumor stroma as a novel treatment strategy for breast cancer: shifting from the neoplastic cell-centric to a stroma-centric paradigm. Adv Pharmacol. 2012;65:45–61. https://doi.org/10.1016/B978-0-12-397927-8.00003-8.
72. West HJ. Can we define and reach precise goals for precision medicine in cancer care? J Clin Oncol. 2016;34(30):3595–6. https://doi.org/10.1200/JCO.2016.68.8226.
73. Hesslow G. Do we need a concept of disease? Theor Med. 1999;14(1):1–14. https://doi.org/10.1007/BF00939846.
74. Reznek L. The nature of disease. New York: Routledge; 1987.
75. Kroschinsky F, Stölzel F, von Bonin S, Beutel G, Koczekan M, Kiehl M, Schellongowski P. Intensive Care in Hematological and Oncological Patients (iCHop) Collaborative Group. New drugs, new toxicities: severe side effects of modern targeted and immunotherapy of cancer and their management. Crit Care. 2017;21(1):89. https://doi.org/10.1186/s13054-017-1678-1.
76. Prasad V. Our best weapons against cancer are not magic bullets. Nature. 2020;577(7791):451. https://doi.org/10.1038/s41586-020-00116-2.
77. Porter R. The greatest benefit to mankind. A medical history of humanity from antiquity to the present. London: Harper Collins; 1997.
78. Stefl S, Nishi H, Petukh M, Panchenko AR, Alexov E. Molecular mechanisms of disease-causing missense mutations. J Mol Biol. 2013;425(21):3919–36. https://doi.org/10.1016/j.jmb.2013.07.014.
79. Wade DT, Halligan PW. Do biomedical models of illness make for good healthcare systems? BMJ. 2004;329(7479):1398–401. https://doi.org/10.1136/bmj.329.7479.1398.
80. Smith R. In search of “non-disease.” BMJ. 2020;372(3742):883–5. https://doi.org/10.1136/bmj.m372.883.
81. Scannell JW, Blanckley A, Boldon H, Warrington B. Diagnosing the decline in pharmaceutical R&D efficiency. Nat Rev Drug Discov. 2012;11(3):191–200. https://doi.org/10.1038/nrd3681.
82. Kola I, Bell J. A call to reform the taxonomy of human disease. Nat Rev Drug Discov. 2011;10(9):641–2. https://doi.org/10.1038/nrd3534.
83. Thorley-Lawson DA. Basic virological aspects of Epstein-Barr virus infection. Semin Hematol. 1988;25(3):247–60.
84. Wright DH. (1978) Burkitt’s lymphoma and infectious mononucleosis. In: The Immunopathology of Lymphoreticular Neoplasms. J.J. Twomey and R.A. Good (Ed.) New York: Plenum Pub. Corp.
85. Giller RH, Grose C. Epstein-Barr virus: the hematologic and oncologic consequences of virus-host interaction. Crit Rev Oncol Hematol. 1989;9(2):149–95. https://doi.org/10.1016/s1040-8428(99)80009-5.
86. Grömminger S, Mautner J, Bornkamm GW. Burkitt lymphoma: the role of Epstein-Barr virus revisited. Br J Haematol. 2012;156(6):719–29. https://doi.org/10.1111/j.1365-2411.2011. 90007.x.
87. Rezk SA, Zhao X, Weiss LM. Epstein-Barr virus (EBV)-associated lymphoid proliferations, a 2018 update. Hum Pathol. 2018;79:18–41. https://doi.org/10.1016/j.humpath.2018.05.020.
88. Naseem M, Barzi A, Brezden-Masley C, Puccini A, Berger MD, Mastria G, Mancini V, Viganò A, Di Piero V. Alice in wonderland syndrome: a clinical and pathophysiological review. Biomed Res Int. 2016. https://doi.org/10.1155/2016/8243145.
89. Ascherio A, Munger KL, EBV and autoimmunity. Curr Top Microbiol Immunol. 2015;390(Pt 1):365–85. https://doi.org/10.1007/978-3-319-22822-8_15.
90. Nussinovitch M, Prais D, Volovitz B, Shapiro R, Amir J. Post-infectious acute cerebellar ataxia in children. Clin Pediatr (Phila). 2003;42(7):581–4. https://doi.org/10.1177/000990020304200720.
91. Plattner R, Anderson MJ, Sato KY, Fasching CL, Der CJ, Stanton BF. Post-infectious acute cerebellar ataxia in children. Proc Natl Acad Sci U S A. 1996;93(13):6665–70. https://doi.org/10.1073/pnas.93.13.6665.
92. Barker DJ. The developmental origins of adult disease. J Am Coll Nutr. 2004. https://doi.org/10.1080/07315724.2004.10719428.
93. Barker DJ. The fetal and infant origins of adult disease. BMJ. 1990;301(6761):1111. https://doi.org/10.1136/bmj.301.6761.1111.
94. Soto AM, Sonnenschein C. Endocrine disruptors - putting the mechanistic cart before the phenomenological horse. Nat Rev Endocrinol. 2018;14(6):317–8. https://doi.org/10.1038/s41574-018-0003-7.
95. Speroni L, Voutilainen M, Mikkola ML, Klager SA, Schaerbecher CM, Sonnenschein C, Soto AM. New insights into fetal mammary gland morphogenesis: differential effects of natural and estrogenic environmental exposures. Sci Rep. 2017. https://doi.org/10.1038/srep40806.
96. Soto AM, Brisken C, Schaerbecher C, Sonnenschein C. Does cancer start in the womb? altered mammary gland development and predisposition to breast cancer due to in utero exposure to endocrine disruptors. J Mammary Gland Biol Neoplasia. 2013;18(2):199–208. https://doi.org/10.1007/s10911-013-9293-5.
97. Zambrowicz BP, Sands AT. Modeling drug action in the mouse with knockouts and RNA interference. Drug Discov Today Targets. 2004;3:198–207.
98. Barabási AL, Oltvai ZN. Network biology: understanding the cell’s functional organization. Nat Rev Genet. 2004;5(2):101–13. https://doi.org/10.1038/nrg1272.
99. Wagner A, Wright J. Alternative routes and mutational robustness in complex regulatory networks. Biosystems. 2007;88(1–2):163–72. https://doi.org/10.1016/j.biosystems.2006.06.002.
100. Rychkov GN, Ilatovskiy AV, Nazarov IB, Shvetsov AV, Lebedev DV, Konev AV, Isaev-Ivanov VV, Onufriev AV. Partially assembled nucleosome structures at atomic detail. Biophys J. 2017;112(3):460–72. https://doi.org/10.1016/j.bpj.2016.10.041.
101. Mojtahedi M, Skupin A, Zhou J, Castaño IG, Leong-Quong RY, Chang H, Trachana K, Giuliani A, Huang S. Cell fate decision as high-dimensional critical state transition. PLoS Biol. 2016. https://doi.org/10.1371/journal.pbio.2000640.
102. Bizzarri M, Giuliani A, Minini M, Monti N, Cucina A. Constraints shape cell function and morphology by canalizing the developmental path along the Waddington’s landscape. bioessays. 2020. https://doi.org/10.1002/bies.201900108.
103. Venegas JG, Winkler T, Musch G, Vidal Melo MF, Layfield D, Tgavealekos N, Fischman AJ, Callahan RJ, Bellani G, Harris RS. Self-organized patchiness in asthma as a prelude to catastrophic shifts. Nature. 2005;434(7034):777–82. https://doi.org/10.1038/ nature03490.
104. Csermely P, Korcsmáros T, Kiss HJ, London G, Nussinov R. Multicomponent therapies for networked systems. Nat Rev Drug Discov. 2005;4(1):71–8. https://doi.org/10.1038/nrd1272.
105. Keith CT, Borsy AA, Stockwell BR. Phenotypic plasticity and the origins of diversification: a network biomarkers. PLoS Comput Biol. 2017. https://doi.org/10.1371/journal.pcbi.1005633.
106. Eriksson JG, Forsén T, Tuomilehto J, Osmond C, Barker DJ. Early growth and coronary heart disease in later life: longitudinal study. BMJ. 2001;322(7292):949–53. https://doi.org/10.1136/bmj.322.7292.949.
107. Lithell HO, McKeigue PM, Berglund L, Mohsen R, Lithell UB, Leon DA. Relation of size at birth to non-insulin dependent diabetes and insulin concentrations in men aged 50–60 years. BMJ. 1996;312(7028):406–10. https://doi.org/10.1136/bmj.312.7028.406.
108. West-Eberhard MJ. Phenotypic plasticity and the origins of diversity. Ann Rev Ecol Syst. 1989;20:249–78.
109. Waddington C. Canalization of development and the inheritance of acquired characters. Nature. 1942;150:563–5.
110. Liu X, Chang X, Liu R, Yu X, Chen L, Aihara K. Quantifying critical states of complex diseases using single-sample dynamic network biomarkers. PLoS Comput Biol. 2017. https://doi.org/10.1371/journal.pcbi.1005633.
111. Chen L, Liu R, Liu ZP, Li M, Aihara K. Detecting early-warning signals for sudden deterioration of complex diseases by dynamical network biomarkers. Sci Rep. 2012. https://doi.org/10.1038/srep00342.
112. Juengst ET, Settersten RA Jr, Fishman JR, McGowan ML. After the revolution? Ethical and social challenges in “personalized genomic medicine.” Per Med. 2012;9(4):429–39. https://doi.org/10.2217/pme.12.37.
113. Evans BJ, Burke W, Jarvik GP. The FDA and genomic tests—getting regulation right. N Engl J Med. 2015;372(23):2258–64. https://doi.org/10.1056/NEJMrs1501194.
114. Pilié PG, LoRusso PM, Yap TA. Precision medicine: progress, pitfalls, and promises. Mol Cancer Ther. 2017;16(12):2641–4. https://doi.org/10.1158/1535-7163.MCT-17-0904.
115. Baird P. The Human Genome Project, genetics and health. Community Genet. 2001;4(2):77–80. https://doi.org/10.1158/0005 1161.
120. Schleidgen S, Klingler C, Bertram T, Rogowski WH, Marckmann G. What is personalized medicine: sharpening a vague term based on a systematic literature review. BMC Med Ethics. 2013;14:55. https://doi.org/10.1186/1472-6939-14-55.

121. Covid-19 Raises Questions About the Value of Personalized Medicine. https://www.wsj.com/articles/covid-19-raises-questions-about-the-value-of-personalized-medicine-1158894927

122. Noble D. A theory of biological relativity: no privileged level of causation. Interface Focus. 2012;2(1):55–64. https://doi.org/10.1098/rsfs.2011.0067.

123. Skladany L, Koller T, Adamcova Selcanova S, Vnencakova J, Jancekova D, Durajova V, Laffers L, Svac J, Janickova K, Palkovic M, Kohout P, Golubnitschaja O. Challenging management of severe chronic disorders in acute pandemic situation: chronic liver disease under COVID-19 pandemic as the proof-of-principle model to orchestrate the measures in 3PM context. EPMA J. 2021;12(1):1–14. https://doi.org/10.1007/s13167-021-00231-8.

124. Wheler JJ, Janku F, Naing A, Li Y, Stephen B, Zinner R, Subbiah V, Fu S, Karp D, Falchook GS, Tsimberidou AM, Piha-Paul S, Anderson R, Ke D, Miller V, Yelensky R, Lee JJ, Hong DS, Kurock R. Cancer therapy directed by comprehensive genomic profiling: a single center study. Cancer Res. 2016;76(13):3690–701. https://doi.org/10.1158/0008-5472.CAN-15-3043.

125. Moscow JA, Fojo T, Schilsky RL. The evidence framework for best survival outcomes in palliative treatment of liver malignancies: unsupervised machine learning and 3 PM recommendations. EPMA J. 2020;11(3):505–15. https://doi.org/10.1007/s13167-020-00221-2.

126. Skladany L, Koller T, Adamcova Selcanova S, Vnencakova J, Jancekova D, Durajova V, Laffers L, Svac J, Janickova K, Palkovic M, Kohout P, Golubnitschaja O. Challenging management of severe chronic disorders in acute pandemic situation: chronic liver disease under COVID-19 pandemic as the proof-of-principle model to orchestrate the measures in 3PM context. EPMA J. 2021;12(1):1–14. https://doi.org/10.1007/s13167-021-00231-8.

127. Wheler JJ, Janku F, Naing A, Li Y, Stephen B, Zinner R, Subbiah V, Fu S, Karp D, Falchook GS, Tsimberidou AM, Piha-Paul S, Anderson R, Ke D, Miller V, Yelensky R, Lee JJ, Hong DS, Kurock R. Cancer therapy directed by comprehensive genomic profiling: a single center study. Cancer Res. 2016;76(13):3690–701. https://doi.org/10.1158/0008-5472.CAN-15-3043.

128. Gerner C, Costigliola V, Golubnitschaja O. Multiomic patterns associated pathologies, prediction, prevention and personalisation of transcript and metabolic profiles: a new approach in systems biology. EMBO Rep. 2003;4(10):989–93. https://doi.org/10.1002/embor.embor944.

129. Golubnitschaja O, Polivka J, Yeghiazaryan K, Berliner L. Liquid biopsy and multiparametric analysis in management of liver malignancies: new concepts of the patient stratification and prognostic approach. EPMA J. 2018;9(3):271–85. https://doi.org/10.1007/s13167-018-0146-6.

130. Bizzarri M, Pontecorvi P. Critical transition across the Waddington landscape as an interpretative model: comment on “Dynamic and thermodynamic models of adaptation” by A.N. Gorban et al. Phys Life Rev. 2021;38:115–9. https://doi.org/10.1016/j.phyrev.2021.05.010.0.

131. Kucera R, Pecen L, Topolcan O, Dahal AR, Costigliola V, Giordano FA, Golubnitschaja O. Prostate cancer management: chronic liver disease under COVID-19 pandemic as the proof-of-principle model to orchestrate the measures in 3PM context. EPMA J. 2021;12(1):1–14. https://doi.org/10.1007/s13167-021-00231-8.

132. Lu M, Zhan X. The crucial role of multiomic approach in cancer research and clinically relevant outcomes. EPMA J. 2018;9(1):77–102. https://doi.org/10.1007/s13167-018-0128-8.

133. Naranjo JD, Saldin LT, Sobieski E, Quijano LM, Hill RC, Chan PG, Torres C, Dzikzi JL, Cramer MC, Lee YC, Das R, Bajwa AK, Nossair R, Klimak M, Marchal L, Patel S, Velankar SS, Hansen KC, McGrath K, Badylak SF. Esophageal extracellular matrix hydrogel mitigates metaplastic change in a dog model of Barrett’s esophagus. Sci Adv. 2020;6(27):eaba4526. https://doi.org/10.1126/sciadv.aba4526.

134. Bizzarri M, Pontecorvi P. Critical transition across the Waddington landscape as an interpretative model: comment on “Dynamic and thermodynamic models of adaptation” by A.N. Gorban et al. Phys Life Rev. 2021;38:115–9. https://doi.org/10.1016/j.phyrev.2021.05.010.0.

135. Proietti S, Cucina A, Pensotti A, Biava PM, Minini M, Monti N, Catizone A, Ricci G, Leonetti E, Harrath AH, Alwasel SH, Bizzarri M. Active fraction from embryo fish extracts induces reversion of the malignant invasive phenotype in breast cancer through down-regulation of TCTP and modulation of E-cadherin/β-catenin pathway. Int J Mol Sci. 2019;20(9):2151. https://doi.org/10.3390/ijms20092151.

136. Bizzarri M, Pontecorvi P. Critical transition across the Waddington landscape as an interpretative model: comment on “Dynamic and thermodynamic models of adaptation” by A.N. Gorban et al. Phys Life Rev. 2021;38:115–9. https://doi.org/10.1016/j.phyrev.2021.05.010.0.