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Title

Precision Medicine for CRC Patients in the Veteran Population: State-of-the-Art, Challenges and Research Directions.

Permalink

<https://escholarship.org/uc/item/6gm9c45q>

Journal

Digestive diseases and sciences, 63(5)

ISSN

0163-2116

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Publication Date

2018-05-01


DOI

10.1007/s10620-018-5000-0

Peer reviewed



Precision Medicine for CRC Patients in the Veteran Population: State-of-the-Art, Challenges and Research Directions

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Received: 2 November 2017 / Accepted: 23 February 2018 / Published online: 23 March 2018
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Abstract

Colorectal cancer (CRC) accounts for ~9% of all cancers in the Veteran population, a fact which has focused a great deal of the attention of the VA's research and development efforts. A field-based meeting of CRC experts was convened to discuss both challenges and opportunities in precision medicine for CRC. This group, designated as the VA Colorectal Cancer Cell-genomics Consortium (VA4C), discussed advances in CRC biology, biomarkers, and imaging for early detection and prevention. There was also a discussion of precision treatment involving fluorescence-guided surgery, targeted chemotherapies and immunotherapies, and personalized cancer treatment approaches. The overarching goal was to identify modalities that might ultimately lead to personalized cancer diagnosis and treatment. This review summarizes the findings of this VA field-based meeting, in which much of the current knowledge on CRC prescreening and treatment was discussed. It was concluded that there is a need and an opportunity to identify new targets for both the prevention of CRC and the development of effective therapies for advanced disease. Also, developing methods integrating genomic testing with tumoroid-based clinical drug response might lead to more accurate diagnosis and prognostication and more effective personalized treatment of CRC.

Keywords Biomarkers · Cancer stem cells · Clinical drug response · Colorectal cancer · FiSS · Genomic testing · Next generation sequencing · Precision Oncology Program (POP) · Tumoroids

Introduction

Colorectal cancer (CRC) is the second most common cause of cancer-related deaths in the USA with an estimated 5% lifetime risk [1]. Another 3% of individuals diagnosed with CRC have a high-risk hereditary cancer syndrome such as Lynch syndrome. In recent years, a major emphasis has been placed on screening of conventional adenomatous and serrated polyps with the intention of removing them prior to their progression to CRC. The serrated polyp has been recognized as a progenitor of 15–30% of all CRCs. Technological advancements in CRC screening procedures and modalities, such as fecal immunochemical test (FIT), air contrast barium enema, and colonoscopy, have led to a decrease in mortality from CRC since the mid-1980s [1]. Despite these

screening tools, only 40% of CRCs are diagnosed at an early stage (I or II) [2]. This is due in part to a lack of patient compliance, limited access to screening colonoscopy, and the low sensitivity/specificity of more commonly used tests such as FIT [2]. Furthermore, a “one drug fits all” cancer treatment approach has been unsuccessful in effectively reducing cancer deaths. For these reasons, there is an urgent need to identify new strategies for the prevention of CRC as well as the development of effective therapeutics for advanced disease.

Specifically, the US Veteran population represents 3% of all cancer patients in the USA and CRC accounts for ~9% of all cancers among Veterans [4]. This is because a large proportion of Veterans receiving care under Veterans Health Administration (VHA) have poor health, multiple comorbidities, and lower income [3]. Further, the incidence of cancer among Veterans is likely to increase with the aging population, an increase in military personnel, and an increase in Veterans seeking health care in VHA [4]. VHA has

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supported a Precision Oncology Program (POP) that uses multi-gene, next-generation sequencing (NGS) panels for patients diagnosed with Stage III or IV CRC, with an intent to provide precision cancer treatment (PCTx). These panels screen for ~200 genes, cost about \$1500 per panel, have a turnaround time of about 2 weeks, and are not validated for PCTx use. Also, since genomic testing only provides a predictive approach, several tumor spheroid culture systems have been examined for their potential use in PCTx, but this technology also has several critical barriers. Thus, there is a need to integrate genomics and cell-based approaches, referred to as “cell-genomics,” to accurately diagnose, prognosticate, and treat gastrointestinal malignancies including pancreatic and CRCs. To this end, in a VA R&D supported field-based meeting, a panel of CRC experts met to review research advances on the biology, biomarkers, and treatment of CRC. The panel discussed the state-of-the-art science, challenges, and opportunities and pragmatically established a “VA Colorectal Cancer Cell-genomics Consortium” (VA4C) with the goal of fostering multi-site collaborative studies aimed at early detection, prevention, and diagnosis as well as novel treatments for CRC. This review summarizes these discussions on the latest developments in biology and biomarkers to individualize diagnostics and prognostication and precision oncology approaches, and charts future collaborative research directions.

Precision Diagnostics and Prognostics for CRC

Advances in Biology, Biomarkers, and Imaging for Early Detection of CRCs

To date the principal modalities for identifying early-stage CRCs have been through use of screening colonoscopy or fecal-based methods such as FIT. The rationale for this approach is to identify either polyps (precancerous) or early-stage CRCs before they develop into lethal tumors. Early-stage CRC is curable with surgical treatment, whereas advanced or metastatic disease may only be treated to prolong survival without curative intent [5]. The traditional paradigm is that sporadic CRC progresses in a stepwise fashion from normal epithelium, to aberrant crypt foci, to low-grade then high-grade adenomas and ultimately to adenocarcinoma which may then metastasize to other organs. This pattern of tumor progression was first identified by Vogelstein and colleagues some 30 years ago [6]. It is now known that cancer cell heterogeneity and other factors can modulate this progression. For example, recent studies suggest that CRCs originating from different locations may have different metastatic targets, for example, colon cancer spreads primarily to liver, whereas rectal cancer spreads primarily to lung

[7]. Moreover, early in tumorigenesis, some cells within the primary tumor already possess the ability to invade and seed distant organs [8]. Such cells have acquired mesenchymal traits through the epithelial-to-mesenchymal transition (EMT). EMT is defined by a variety of genetic and morphological changes, which allow cells to invade normal colon epithelium, vascular endothelium, and eventually the parenchyma of target organs. Recent work indicates that the genetic signatures identifying EMT may also be cell context-dependent [9]. Metastatic cell deposits may grow slowly or lie dormant until circulating and stroma-dependent factors stimulate their arousal and growth.

An important anatomical factor very specific to CRC is the recently identified “serrated polyp-neoplasia” pathway [10–12]. Sessile serrated adenoma/polyps (SSA/Ps), established precursors of CRC, exhibit microsatellite instability (MSI) and account for approximately 15–30% of sporadic CRCs [13]. SSA/Ps are predominantly located in the right/proximal colon and observed in “interval cancers” occurring within a short time after a screening procedure (2–6% of all CRCs) [14, 15]. Serrated polyps are characterized by a variety of histologic features including irregular distribution of crypts, dilatation of crypt bases, serration at crypt bases, branched crypts, horizontal extension of crypt bases, dysmaturation of crypts, and herniation of crypts through the muscularis mucosae [16]. According to the American Gastroenterology Association criteria, even one crypt showing any of the above characteristics is sufficient to diagnosis SSA/P. Approximately 17.6% of SSA/Ps (> 2 cm) show residual adenomatous tissues when reexamined [17]. Unlike other adenomatous polyps, SSA/Ps are characterized by mutations in the BRAF proto-oncogene that causes the development of polyps. Hyper-methylation of CpG islands in the promoter regions of tumor suppressor genes, including *MLH1*, leads to a decrease in tumor suppression and an increase in sporadic MSI [10, 18–21]. Because benign hyperplastic polyps (HPs) and SSA/Ps have a different risk of developing into malignant disease, their accurate classification is critical for making clinical decisions (e.g., surveillance intervals, intervention). Understanding these complex processes and testing all prognostic and therapeutic measures in animal models before proceeding to clinical trials is challenging. It is noteworthy that although colonoscopic screening and polypectomy have led to a decline in advanced CRC, its utility for differentiating SSA/Ps from HPs has not yet been realized.

Genetic and Epigenetic CRC Biomarkers

The multistep process of CRC development is associated with the accumulation of genetic and epigenetic alterations in epithelial cells via distinct molecular pathways, including the common “adenoma-carcinoma” pathway as

well as the recently identified “serrated polyp-neoplasia” pathway [6–8]. Hence, identifying these CRC molecular markers, i.e., biomarkers, is key to early detection, staging, and assessment of therapeutic efficacy, as reviewed recently [22]. Whereas current guidelines recommend surgery alone for Stage I and adjuvant chemotherapy for Stage III cancers, the appropriate treatment for Stage II cancers, which have breached the colon wall but not invaded lymph nodes or other organs, is uncertain. Although it is proposed that molecular biomarker panels, including genetic, epigenetic, protein and carbohydrate-based markers, may identify Stage II cancers that warrant aggressive treatment after surgical resection [23], none have been validated prospectively.

Circulating tumor DNA and RNA-based biomarkers offer high specificity and are ideal as predictive biomarkers for monitoring the response to chemotherapy as well as tumor progression, as reviewed recently [24]. However, due to low DNA yields and low mutational burden, they are inadequate for diagnosis. Cell-free microRNAs (miRNAs) and small non-coding RNAs (ncRNAs) are now emerging as noninvasive and highly sensitive biomarkers, but specificity remains a concern. For these reasons, large-scale prospective clinical studies aimed at carefully evaluating the sensitivity and specificity of these biomarkers are needed before their adoption to clinical practice.

Hereditary cancer syndromes have provided powerful insights into our understanding of somatic mutations present in sporadic cancers, as well as implicated cell signaling pathways [20–24]. One clear example is the identification of germline, inactivating mutations in the APC gene, which encodes a 300-kD wnt pathway adaptor protein [23]. Although germline mutations in APC are responsible for familial adenomatous polyposis (FAP), a rare condition affecting about 1 in 7000 individuals in the USA, somatic mutations in the APC gene are present in more than 70% of colonic adenomatous polyps and carcinomas [24]. Hereditary non-polyposis colon cancer (HNPCC), or Lynch syndrome, involves a germline mutation in a DNA mismatch repair (MMR) genes, leading to defective DNA repair. Recent studies identifying inactivation of TGF- β signaling through loss of the Smad3 adaptor β -2 spectrin could potentially provide new insights into CRC development from stem-like tumor-initiating cells (STICs) for targeted chemoprevention.

Further, genome-wide association studies (GWAS) have identified numerous genetic loci that are associated with CRC, but heritability remains unknown. More than 40 low-penetrant polymorphic variants are reported [25]. More recently in a Finnish cohort, a single-nucleotide polymorphism (SNP) rs992157 at chromosome 2q35 intronic to PNKD and TMBIMI was found to be associated with CRC [26]. Genome-wide alterations in DNA methylation influence gene expression resulting in altered aberrant crypt foci,

an early precursor of CRC [27]. Recently, single-cell RNA sequencing (e.g., GemCode technology 10 \times Genomics, San Francisco, CA) has permitted an assessment of transcriptional events that may be important in the development of cancers and could potentially predict therapeutics. In addition to genomic profiling approaches, depending upon the molecule of interest, several additional “omics” approaches are being utilized, including proteomics, metabolomics, lipidomics (chloroform/methanol), glycomics (glycoprotein isolation), and peptidomics.

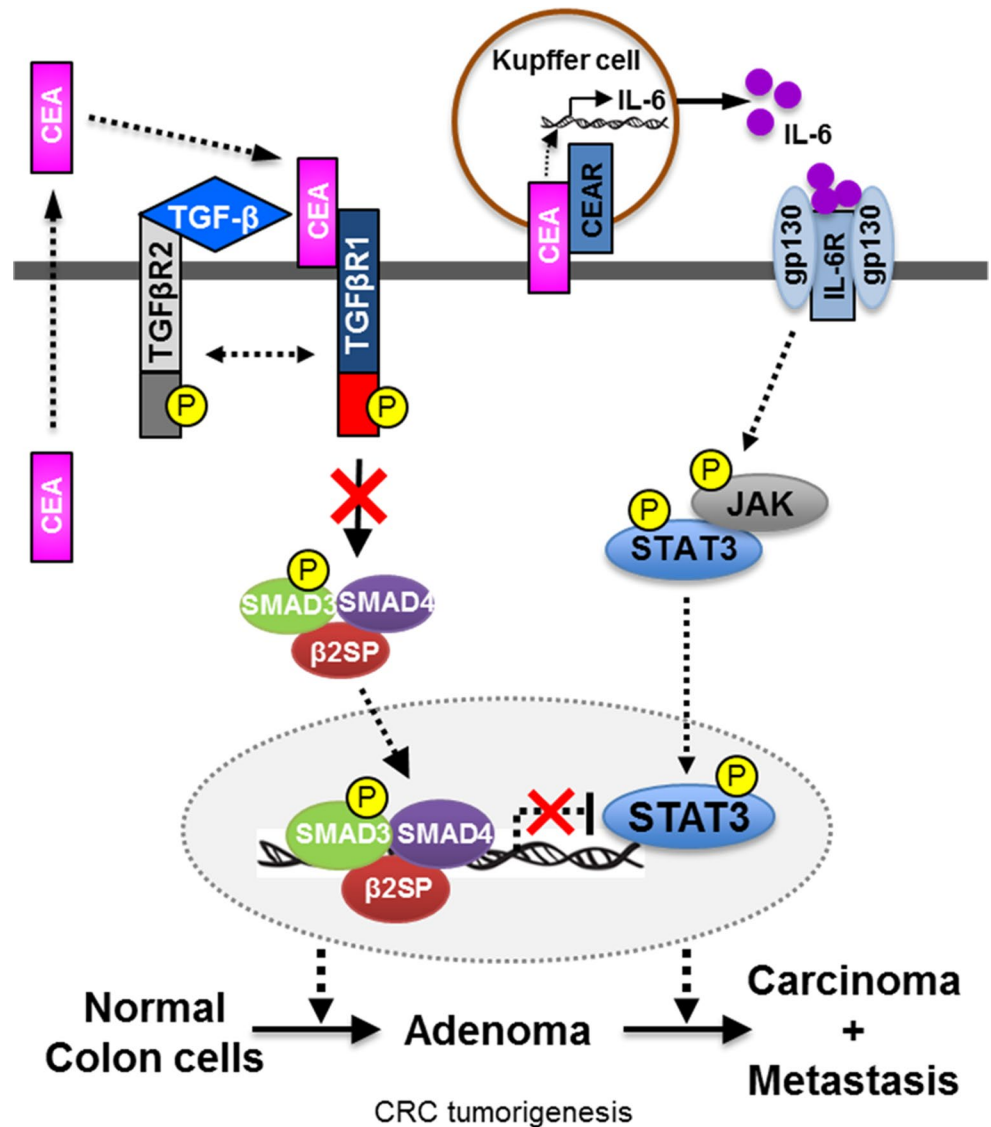
TGF- β Receptor and SMAD Gene Mutations in Gastrointestinal/CRC

In normal and premalignant cells, TGF- β enforces homeostasis and suppresses tumor progression through tumor-suppressive effects on the stroma. Current data strongly support TGF- β signaling as a suppressor of early CRCs [28, 29]. In advanced disease, metastatic CRCs escape the tumor suppressor effects of TGF- β signaling by becoming resistant to TGF- β -induced growth inhibition [30]. Inactivating mutations in the *TGFBR2* gene occur in most human CRC and gastric carcinomas that demonstrate MSI [31]. The TGF- β signaling network is also disrupted in cancer by mutations in *Smad4*, which occur in more than 30% of CRCs [32, 33]. Smad-deficient mice display phenotypes that suggest a tumor suppressor role for the Smads [34]. When mice with one mutated APC allele are crossed with heterozygous Smad4 mice, the compound heterozygotes develop larger polyps that can progress into malignant adenocarcinomas [35]. Although fewer Smad3 mutations have been found in human cancers, mice with a Smad3 homozygous deletion develop aggressive CRC at an early age in a manner that seems to be highly dependent on the genetic background of the mice.

Cross talk between CEA, TGF- β signaling, and STAT3

Figure 1 depicts cross talk between carcinoembryonic antigen (CEA), TGF- β signaling, and STAT3. Advanced CRC is associated with high levels of circulating CEA and a dismal survival; the elevated levels of CEA are especially associated with Stage IV cancers. Consequently, there is a need to identify new targets for both the prevention of CRC as well as the development of novel therapies for advanced cancer. CEA (also known as CEACAM5) is one of the few biomarkers approved by the U.S. Food and Drug Administration (FDA) for CRC [36]. TCGA studies have validated that elevated CEA levels and disruption of TGF- β signaling are observed in over 80% of right-sided CRCs [37]. Activation of STAT3 is also observed in this subgroup of patients. TGF- β induces the secretion of CEA in a dose-dependent manner [38]. Given the importance of CEA, TGF- β , and STAT3 signaling pathways in CRC tumorigenesis, cross talk

Fig. 1 Schematics of cross talk between CEA, TGF- β signaling, and IL6-STAT3 pathway



between these pathways is a focus of investigation. CEA and CEACAM6 are known Smad3 target genes. An important direction for the future is designing studies that delineate the molecular mechanisms underlying the cross talk between CEA and the TGF- β and STAT3 signaling pathways during CRC development.

Mucins as Biomarkers for CRC Early Detection and Prognosis

Several studies showed moderate to high inter-observer disagreement between pathologists in the histologic differentiation of SSA/Ps (highly malignant) and HPs (benign) [21, 39–46]. Considering this variability, studies have focused on identifying molecular markers for distinguishing between these two entities. For example, MUC5AC, TFF1, and Annexin A10 have emerged as potential markers for

differentiating benign polyps from SSA/P [47, 48]. Aberrant changes in the expression, localization, and glycosylation of colonic mucin have been observed in very early stages of colon cancer. SSA/Ps exhibit an elevated expression of trans-membrane mucin MUC17 and secreted mucin MUC5AC in comparison to HP [49]. Notably, in conjunction with these expressional changes, distinct differences were also observed among the studied polyp groups in MUC5AC and MUC4 localization. This signifies the ability of these markers to discriminate benign HP from SSA/P polyp subtypes. In multivariate regression models in conjunction with ROC curve analyses, the combination of MUC17/MUC5AC effectively discriminated SSA/Ps from HP [49]. The trio (CA 19-9, MUC17, and MUC5AC) emerged as a combined panel for discriminating polyp subtypes and thus accurately classifying colorectal polyps. Based on the lack of clear tools to identify SSA/Ps, histologic markers can aid in improving

the differentiation of SSA/PPs from benign polyp subtypes. Due to difficulties in their diagnosis by endoscopists, their inter-observer variability among pathologists, and their highly malignant nature, a means of identifying SSA/PPs is central to the improved detection and timely treatment of CRC. There is a broad consensus that future studies should focus on identifying alternative molecular markers that can more successfully achieve these goals.

Biomarkers for Colon Cancer Dissemination

One novel approach to stratify the malignant potential of CRC is to identify cancer cells with distinct invasive potential at the tumor “front,” namely the interface between tumor and normal tissue. Nuclear accumulation of β -catenin distinguishes the cells at the “front” from those closer to the tumor core [50]. This is consistent with observations that cells tend to invade in clusters, not as individuals [9]; these cells likely share attributes that predispose them to behave as an aggressive unit. Cell clusters at the tumor front are more likely to co-overexpress M3 muscarinic receptors (M3R) and matrix metalloproteinase-1 (MMP1) within cytoplasmic vesicles [51]. This is potentially important since M3R activation robustly induces expression of MMP1, a secreted collagenase that promotes colon cancer invasion [52–54]. Validating the use of such histochemical tests in formalin-fixed paraffin-embedded tissues may yield an efficient and economical way to identify those Stage II cancers that require more aggressive therapy.

Histopathology with Neuroendocrine Tumor (NET) Stains Identifies Neuroendocrine Features of CRC

A subset of adenocarcinomas and adenomas possess neuroendocrine features [55]. Neuroendocrine cells possess bioactive amines and peptides that may have an important role in promoting the growth of transformed cells such as those within adenocarcinomas, as has been demonstrated with peptides

such as PACAP [56]. The expression of receptors on CRCs can differentially couple to intracellular signaling molecules that promote proliferation [57]. A recent study identified neuroendocrine features in up to one-third of 40 patients with CRC (unpublished data); assays of chromogranin A and neuron-specific enolase were performed (Fig. 2). The presence of neuroendocrine features may suggest a stem cell component or portend a more aggressive tumor state. Neuroendocrine features may also suggest different chemotherapeutic or biologic strategies in advanced-stage CRC.

Role of Microbiome in CRCs

Well-established risk factors for colorectal adenomas and cancer include: a personal or family history of CRC or polyps; an inherited condition that predisposes to colon polyps and cancer; inflammatory bowel disease (e.g., Crohn’s disease or ulcerative colitis); obesity; type 2 diabetes; heavy alcohol use; a diet high in red meat and processed meats; smoking; and physical inactivity. Additional risk factors that have been described include fatty liver [58], hepatitis C [59], and Agent Orange exposure [60]; the latter is very relevant to Vietnam Veterans. Modifying known dietary and lifestyle risk factors and chemoprevention with aspirin and nonsteroidal anti-inflammatory drug (NSAID) use are associated with reduced risk for colon polyps and CRC [61]. In relation to the role of the diet and intestinal microbiome in CRC pathogenesis, the microbiome of subjects consuming a Western diet (WD) was compared to that of subjects consuming a high-protein diet (HPD) [62] (Fig. 3a, b). There was a highly significant difference in composition by weighted UniFrac analysis ($p < 10^{-5}$) (Fig. 3b). At the genus level, the HPD microbiome was characterized by expansion of *Akkermansia*, *Ruminococcus*, and *Bacteroides* and depletion of *Lactobacillus* and *Turicibacter* (Fig. 3c). The HPD group had increased abundance of 114 operational taxonomic units (OTUs) selected at 97% similarity (corresponding approximately to

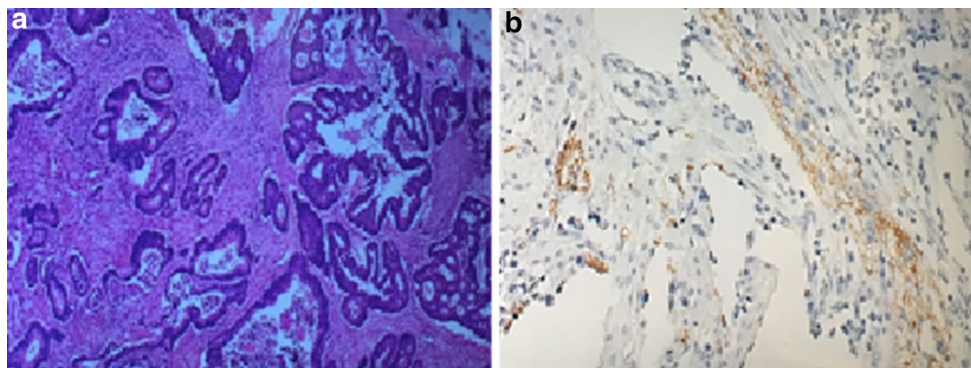


Fig. 2 **a** NET features by H&E staining in colorectal adenocarcinoma specimen. **b** NET features by chromogranin A immunohistochemistry staining in colorectal adenocarcinoma specimen. Unpublished data

species) at a threshold of $q < 0.05$ (representing significance adjusted for multiple hypothesis testing).

Of these, only *Akkermansia mucinophila* ($q = 0.008$) and an unclassified *Clostridiales* ($q = 0.04$) had a statistically significant inverse correlation with fat mass after adjustment for diet [62]. *Fusobacterium* enrichment is associated with specific molecular subsets of CRCs, suggesting that there is a potential pathogenic role in the development of CRC [63]. Furthermore, it has also been shown that environmental pollutants may affect the microbiome, such as was observed with ultrafine particles [64]. Whether ultrafine particles affect the steps involved in colon cancer pathogenesis remains to be investigated.

Precision Imaging for Prevention of CRCs

Screening colonoscopy has emerged as perhaps the most effective lifesaving intervention against CRC to date. Pooled analysis of numerous large observational studies indicates that systematic removal of all visualized polyps including adenomatous and serrated polyps [43, 65] at colonoscopy decreases proximal and distal CRC incidence and mortality by $> 60\%$ [2]. Currently, a VA clinical trial (50,000-subject, prospective randomized controlled study) is underway (Cooperative Studies Program #577) to assess screening colonoscopy versus FIT for reducing CRC deaths. While cancer death rates in persons older than 50 years are now declining, likely due to the adoption of screening programs, recent data indicate that age-specific CRC incidence and mortality are actually rising in those < 50 years of age. Colonoscopy has not been fully protective due to several uncontrollable factors including missed polyps, operator factors [66–69], and an inability to detect atypical [70] serrated polyps and poorly visualized flat neoplasms [71]. Thus, colonoscopy can benefit from assistive technologies that improve the neoplastic polyp detection rate as

a primary performance benchmark [72]. To this end, standard colonoscopy has undergone recent enhancements to improve detection of more subtle lesions and has been supplemented with other technologies that provide “real-time histology” to enable selective removal [73–75]. Clinical trials, however, have shown that both high definition colonoscopy and dye spray chromoendoscopy only marginally improve neoplastic polyp detection [76, 77]. Using retrograde and ultra-wide angle camera systems that reveal surface area between folds can improve neoplastic polyp detection by 23–70% [78, 79]. Selective wavelength imaging and transparent cap-assisted colonoscopy have had mixed results [80, 81]. Thus, despite some improvements there remains a major unmet need in the ability to detect and remove atypical and serrated polyps.

CRC: Advances in Precision Surgery and Treatment

Although surgical resection is the mainstay of treatment in early-stage CRC, incomplete removal of tumors leads to local or widely metastatic disease which is resistant to conventional chemotherapeutics. For patients whose disease has progressed to metastatic CRC, molecular profiling may shed light on why various treatments fail. The discussions in this section focus first on advancing CRC precision treatment through increased precision in CRC surgery. Secondly it reviews the state of the art in targeted chemotherapies.

Fluorescence-Guided Surgery for CRC

Surgical resection for CRC has the greatest potential for cure. Since the application of complete mesocolic excision for colon cancer surgery, local 5-year recurrence rates have decreased [82]. Furthermore, achieving negative

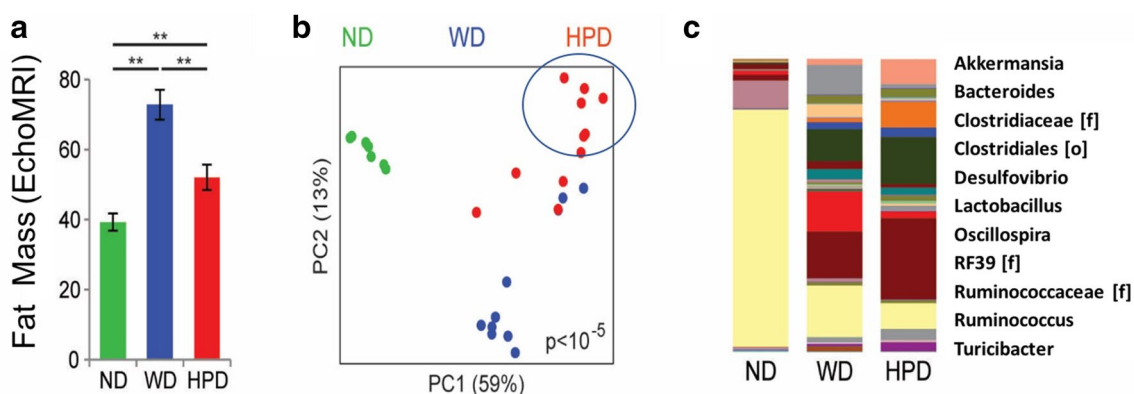


Fig. 3 A high-protein diet induces fat loss and an altered intestinal microbiome in rats with Western diet-induced obesity. **a** Body fat mass of rats kept on a WD or switched to a HPD. Age-matched control rats on a ND are shown for comparison. $**p < 0.01$ **b** Principal

coordinates analysis (PCoA) of the cecal luminal microbiome colored by diet. P value across groups calculated using Adonis. **c** Mean abundance of common genera. Some reads were identified only at the family (f) or order (o) level

microscopic margins and complete resection of metastatic tumor (R0 resection) can significantly improve 5-year survival rates [83–87]. Despite achieving R0 resection rates in patients with CRC, local and distant recurrence has still been reported to be as high as 34% [84, 88, 89]. Furthermore, the prognosis of patients with local recurrence is poor [90]. Real-time, reliable imaging to detect positive surgical margins at the time of surgery would improve outcomes. Emerging techniques are revolutionizing the performance of cancer surgery [91]. Intraoperative fluorescence imaging, or fluorescence-guided surgery (FGS), can provide high fidelity tumor visualization for localization, resection, and margin confirmation [92]. Thus, targeted fluorescent labeling of cancer cells may change the way we find and treat cancer [93, 94].

FGS combines advanced imaging platforms with targeted fluorescent agents to improve their intraoperative detection [95]. One technique is to covalently bind organic dye to a monoclonal antibody which labels a known tumor-specific antigen [91]. One such tumor biomarker is anti-CEA antibodies which label human CRC in murine models [92, 96–98] (Fig. 4). Mouse and chimeric (mouse/human) CEA antibodies, conjugated to fluorescent dyes, are capable of enhancing visualization of submillimeter tumor deposits [92] and aiding with FGS [99–101]. In an effort to improve therapeutic efficacy in the clinic, Yazaki et al. [102] humanized the murine anti-CEA T84.66 antibody with structurally similar “human” segments through a technique known as complementary determining region (CDR) grafting. This humanized anti-CEA antibody is currently used in clinical trials for PET imaging of CRC. In Europe, this technique was recently applied in clinical trials to localize CRC using SGM-101, an antibody-dye conjugate in which the fluorochrome BM104 is coupled to a chimeric monoclonal antibody against CEA [103]. Other fluorophore-conjugated antibodies to tumor-specific antigens such as insulin-like growth factor-1 receptor have been used in preclinical FGS

for colon cancer [104]. In addition, viral vectors such as adenovirus and herpes virus can be used to deliver fluorescent proteins to cancer cells [105]. Activatable cell-penetrating peptides have been engineered to highlight tumors based on enzymatic cleavage of peptidases [106]. Indocyanine green (ICG), used clinically to define liver tumor margins, tissue perfusion, and biliary anatomy, has been shown to reduce leak rates after low anterior resection for CRC [107]. Thus, the use of FGS as a surgical guide has the potential to increase both survival and quality of life for patients [108]. FGS has great potential for a broad range of clinical applications, including in CRC.

Targeted Chemotherapies

Targeted therapies have recently been developed to join traditional therapeutic approaches to CRC including surgery, chemotherapy, and radiotherapy [109]. Targeted chemotherapy may be used for neoadjuvant treatment or adjuvant treatment. For advanced cancers that have spread to other organs, such as the liver, chemo can also be used to help shrink tumors and relieve symptoms. Although it is not likely to cure the cancer, it often prolongs life. Common drugs used for CRC include: 5-fluorouracil (5-FU), often given with leucovorin (also called folinic acid) or levo-leucovorin; capecitabine (Xeloda tablets) metabolized to 5-FU in tumors [110]; irinotecan (Camptosar); and a combination tablet containing oxaliplatin (Eloxatin) and trifluridine and tipiracil (Lonsurf).

About 85% of all sporadic CRCs occur due to chromosomal instability, and ~15% of all CRCs are caused by various deficiencies in the DNA mismatch repair system. In the order of decreasing frequencies, TP53, KRAS, PIK3CA, BRAF, and PTEN are among the most commonly altered genes in CRC [111]. TP53 was the most commonly mutated gene, affecting 17/24 cell lines; hyperactivating KRAS mutations were found in 15 cancer cell lines of which five were

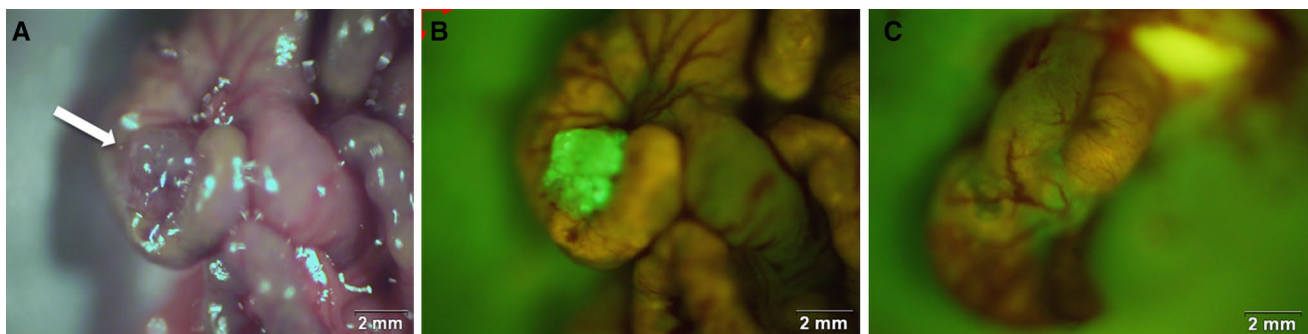


Fig. 4 Labeling of orthotopic human colon tumors in nude mice with fluorophore-conjugated chimeric anti-CEA antibodies. Panel A shows bright-field image of an orthotopic colon tumor (white arrow) and panel B shows the bright fluorescence of the tumor 48 h after labeling

the fluorophore chimeric anti-CEA antibodies. The absence of any fluorescence signal after fluorescence-guided surgery in panel C indicates a complete resection

homozygous, and BRAF mutations were found in another five cell lines.

Several cutting-edge biological modalities are under investigation. Mannosylated liposomes have been used to improve cell uptake and tumor inhibition rate [112]. This was shown without increase in toxicity. Thus, Patras et al. [113] have shown the effects of liposomal paclitaxel encapsulated in long-circulating liposomes together with liposomal prednisolone phosphate to be superior in anti-tumor activity mainly related to anti-angiogenic and anti-inflammatory effects. Attempts have been made to detail the functional pathways microRNA array, and considerable knowledge has been accumulated [114]. Yao et al. [115] have shown the importance of microRNA-215 as a tumor suppressor in colon cancer, which paradoxically acts as oncogene in breast cancer, and have proposed this to be a novel therapeutic target. However, knowledge of several microRNAs is very limited in terms of their targeted genes [113, 114]. Zheng et al. studied the effects of caudal type homeobox (CDX2) and human telomerase reverse transcriptase (hTERT) [115]. hTERT and loss of CDX2 have been associated with colon cancer in humans. In their elegant experiment, they showed over-expression of CDX2 is associated with suppression of colon cancer progression [116]. Finally, as stated before, TGF- β has also been implicated in treatment of metastatic colon cancer [117].

Immunotherapies for CRC

Immune surveillance is a key mechanism to protect against malignancy. Successful CRC immune surveillance relies on the effective function of antigen-presenting cells and T cells. It is now evident that this inadequate function of host immune system is due to suppressive factors like myeloid-derived suppressor cells (MDSCs). It is one of the major mechanisms of tumor escape from immune control as well as an important factor limiting the success of cancer immunotherapy. MDSCs play an important role in tumor non-responsiveness by suppressing antigen-specific T cell responses [118–121]. These cells take up antigen delivered by vaccination, present it to activated T cells, and thereby inhibit the same antigen-specific T cells that the vaccination strategy is aiming to activate. Cancer patients or tumor-bearing mice have a considerable number of MDSCs, which cause even the most effective antigen-delivery strategies to fail [120, 121]. CRC therapy will only be successful if strategies involve either the differentiation or elimination of suppressive cells. This strategy will be more effective than direct cancer therapy that attempts to eliminate tumor cells. Consideration of immune suppressive parameters will provide better stratification of CRC patient treatment.

Currently there are about 46 active (recruiting) immunotherapy trials against CRC (Clin Trials. Gov accessed

on July 23, 2017). Peptide vaccines targeting HER-2/Neu, P-53, SART-3, and CEA in CRC have been studied. These vaccines are generally well tolerated but are far from a successful treatment. Most of the current CRC immunotherapies can be grouped as checkpoint blockade inhibitors like anti-CTLA4 [122], PDL-L1 [123], PD-1 [124], LAG-3; approved FDA targeted antibodies like bevacizumab, cetuximab, ramucirumab, and panitumumab; ACT (adoptive T cell therapy) and virus-mediated therapies. These therapies are tried either as adjuvants or as combinations. Current CRC immunotherapy targets the highly expressed tumor antigens like CEA, APC, MAGE, MUC-1, and survivin [125], or common mutations in the WNT, MAPK, TGF, P53, or apoptosis pathway [126–128]. A recent paper targeting mutant KRAS in CRC patients using ACT observed objective progressive regression [129]. ACT targeting various known CRC mutations in combination with checkpoint inhibitors and modulating the persistent immune suppressive factors would lead to effective therapies. Data from a phase 2 trial of pembrolizumab for the treatment of tumors with and without mismatch repair deficiency support the hypothesis that mismatch repair-deficient tumors are more responsive to PD-1 blockade than are mismatch repair-proficient tumors [130].

Emerging Approaches to Personalizing Cancer Treatment

Genomic Testing

Personalizing cancer treatment that is tailored to an individual's genomic profile is a cornerstone of precision oncology [131]. NGS of DNA from tumor tissue can identify gene variants that could be targets of specific cancer therapies. These variants can be inherited (germline) or acquired (somatic) [132]. For example, poly(ADP-ribose) polymerase (PARP) inhibitors are effective in killing cancer cells that are deficient in repair of double-stranded DNA breaks. This is due to a pathogenic variant in the *BRCA1* or *BRCA2* gene [133] and other germline variants, which cause hereditary breast-ovarian cancer syndrome [134]. Tumors deficient in DNA mismatch repair (MMR) due to a pathogenic variant in the *MSH2*, *MLH1*, *MSH6*, *PMS2*, or *EPCAM* genes are responsive to antibodies or ligands that block the programmed death 1 (PD1) pathway [130]. These germline variants cause Lynch syndrome, the most common cause of hereditary CRC [135].

The VHA has undertaken feasibility studies on the use of NGS testing of tumor samples [136]. Tumors tested with multi-gene NGS sequencing through 1 of 2 contracted vendors were identified from POP records, and cancer characteristics were extracted from POP and medical records. Drug use data were obtained from the VA Corporate Data

Warehouse. NGS testing results and annotations were extracted from POP records. Of a total of 1292 tumor samples sent for NGS testing since program inception in 2015, the most common diagnoses have been lung (561: adeno 418, squamous 143), gastrointestinal [109], lymph node [75], liver [56], head and neck [52], and prostate [43]. The rate of sample test requests increased rapidly after national dissemination in July 2016 (from a mean of 23 samples/month prior to launch to a mean of 126 samples/month 3 months later), as did the number of participating facilities (from a mean of 8/month to a mean of 27/month). Sequencing success rate increased from 68 to 71% over the same interval, while mean turnaround time remained similar at 19.7 and 19.1 days, respectively. NGS was requested for 61 patients with CRC, with almost all requested after dissemination of POP across the USA. Only one case failed to generate NGS results (98% success rate). Twenty-two patients had mutations targetable with an off-label FDA-approved drug, but no patient received NGS-directed treatment. These results suggest that tumor NGS testing as part of POP in the VA healthcare system is feasible with a high rate of testing success in CRC, but effective implementation strategies are needed to ensure uptake of treatments targeting tumor sequencing results.

The Spheroid and Organoid Culture Approaches to Assess Clinical Drug Response

Animal models of human CRC do not recapitulate human disease and frequently yield overly optimistic results [137, 138]. In the absence of accurate murine models, robust cellular models of CRC are considered critical. With over 150 CRC cell lines having been investigated, there is an abundance of cell line resources to investigate [139]. A number of different colon two-dimensional (2D) cell techniques, also referred to as “flat” cell cultures, have been described including the flow cell used for organ-on-chip models [140]. Despite progress made, these flat cultures are significantly different from in vivo tumor cells biochemically, physiologically, and genetically. Another approach that has been developed is in vitro multi-cell tumor-like cultures referred to as spheroids, tumoroids, or organoid culture systems [141]. Whether they are scaffold-free or scaffold-based 3D platforms, they fail to fully recapitulate in vivo tumors [141]. More importantly, they are cumbersome, costly, and not adaptable to perfusion-driven culture. Though very few of these have progressed to clinical testing realm the spheroid technologies are under intense investigation for their use in clinical drug response analyses.

Notably, several studies reported success in predicting CRC clinical drug response using organoid culture, also referred to as histoculture technique. Thus, an evaluation of 86 chemotherapy regimens using the 3D histoculture drug

response assay (HDRA) for CRCs showed that the correlation of HDRA to the clinical response to chemotherapy was 66% (sensitivity 73%, specificity 55%) [142]. In another study, ATP-CRA-guided chemotherapy group showed better treatment response (48.4 vs. 21.9%, $p=0.027$) and a higher rate of resection of hepatic lesions (35.5 vs. 12.5%, $p=0$) [143]. Also, the extreme drug response assay (EDRA), developed as an exclusion test to identify drugs unlikely to elicit a response [144], was used to analyze the therapeutic efficacy of treatment for CRC patients. Thus, in the clinical correlation of 25 Duke’s D patients with EDRA, the sensitivity and specificity of the assay were 100 and 95%, respectively, suggesting that EDRA obtained at initial diagnosis may be useful for the selection of therapeutic regimens for metastatic disease [145, 146]. While these studies indicate the value of ex vivo organoid cultures, these results need to be replicated in multicenter clinical trials.

Tumoroid Technology for Clinical Drug Response

A fiber-inspired smart scaffold (FiSS) tumoroid culture platform has been developed that allows the formation of tumor-like organoids with molecular signatures of tumors. These cancer cells obtained from ex vivo tumor biopsies mimic patient tumors [141, 147, 148] (Fig. 5). The novelty of this platform lies in its simplicity, reproducibility, and similarity to in vivo tumors in terms of its drug responsiveness. Recent pilot proof-of-concept studies showed that cells of tumors from cancer patients, when cultured on a tumoroid platform, formed tumoroids that exhibited differential drug response [141, 147, 148]. Tumoroids were also found to expand cancer stem cells (CSCs) up to 30-fold, with cancer cells originating from several cancer cell lines including cells from breast, lung, and colon tumor xenografts.

In the post-genomic era the development of microarray technology has allowed use of genomic data to help define which patients would benefit most from a specific therapy [149]. For example, gene signatures have been developed and validated against large retrospective databases to risk stratify patients to selectively receive adjuvant treatments for breast cancer [150]. However, genomic/genetic tests are based on pathway analysis of several genes with mutations out of more than 1000 genes that play a role in cancer pathogenesis. They may reveal potential adverse responses to drugs. However, the results do not indicate one or two potential drugs for effective treatments but rather suggest excluding therapies that may cause adverse effects.

One novel approach to PCT involves use of the patient-derived perfused tumoroid (PPT) platform, which mimics patient’s tumor. PPT provides a simple, rapid, scalable, and inexpensive in vitro tumor model that better replicates the structure, physiology, and function of tissues. It also recreates the in vivo morphology and arrangement of individual

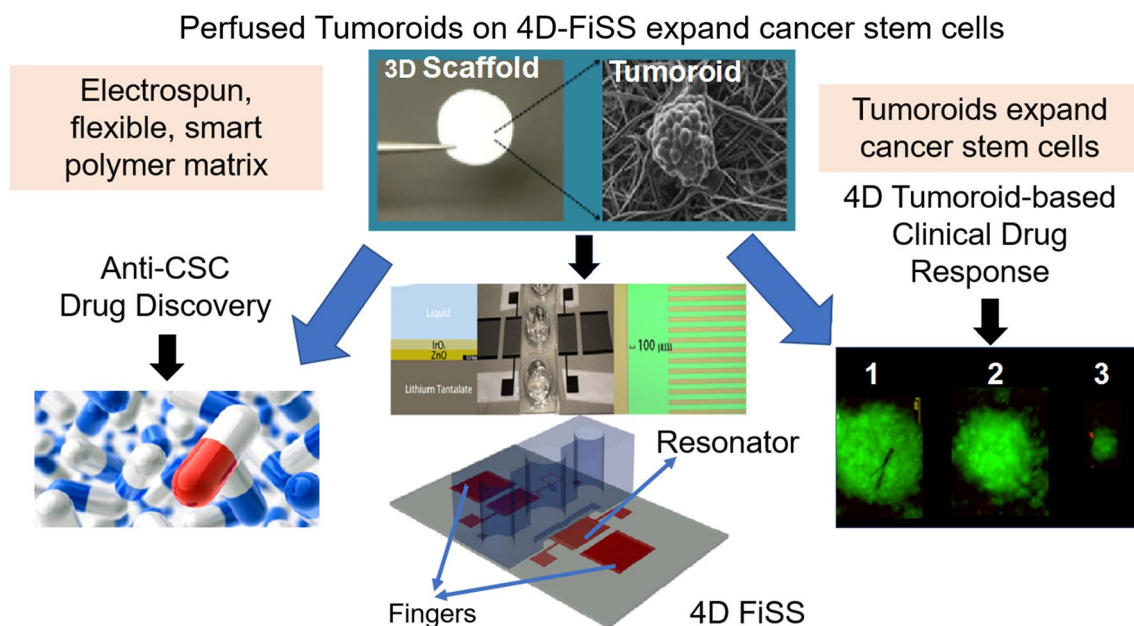


Fig. 5 Schematics of tumoroid technology platform for anti-CSC drug discovery and tumoroid-based clinical drug response assay

cells and the concentration gradients of both signaling molecules and therapeutic agents [151]. This platform can be used to culture tumoroids utilizing human tumor cells (Fig. 5) with or without stromal cells. These are derived from biopsy or surgery discard tumor samples from patients. The PPT also promotes tumoroids in a manner that can be adapted for cell viability measurements (e.g., Celltiter-Glo or PrestoBlue). Preliminary data suggest that PCT tests will fully recapitulate *in vivo* tumors and will provide evidence for the best drug to treat each patient's tumor. Further, expanding PCT to include first genetic/genomic testing that predicts potential therapies, followed by tumoroid-based clinical drug response assay, is expected to significantly improve the standard of personalized therapy. Adding evidence-based, simple-to-analyze, and ready-to-prescribe therapies will significantly improve personalized cancer treatment. Thus, a cell-genomic approach that combines NGS-based genomic testing with a PPT-based clinical drug response assay is expected to provide an exciting approach for PCT and may aid in discovery of novel anticancer agents.

Cancer Stem Cell Biology and Drug Discovery

Pivotal to successful CRC treatments are CSCs, referred to as the “seeds” of tumors, which play a central role in processes such as tumor initiation, drug resistance, invasion, and metastasis. While chemo-, radio- and immune-therapies are more effective at killing bulk tumor cells, the quiescent CSCs manage to escape and seed new tumor growth. CSC phenotype in human colon cancer is associated with

poor prognosis [152, 153]. Hence, a therapeutic approach has been to target CSCs through either inhibition of self-renewal or induction of differentiation [154, 155]. A major barrier to CSC research, however, is the small proportion of these cells in tumors. With the possibility to expand CSCs by tumoroid cultures has opened new avenues for anti-CSC drug discovery. An NCI library of FDA-approved drugs was screened using FiSS tumoroid platform, which revealed that mithramycin A, idarubicin, and daunorubicin were the most potent inhibitors of HT-29 cell viability; and cisplatin, 6-thioguanine, and cytarabine were the least potent inhibitors. Further studies have revealed that mithramycin A treatment can reduce CSC self-renewal as well as expression of stemness genes. These results suggest that mithramycin A may be used as a single agent or in combination with other chemo- or immuno-therapy to most effectively treat or possibly ‘cure’ CRCs.

In addition to these efforts, two major pathways have been exploited to screen for anti-CSC drugs. First, application of HTS on a library of small molecules targeting EMT in breast cancer cells led to identification of four candidates: salinomycin, a polyketide synthase-derived natural product, and probes ML239, ML243, and ML245, which are chemically synthesizable, small hydrophobic molecules [156–160]. A second target signaling pathway for CSCs involves interaction of glycosaminoglycans (GAGs) with factors involved in growth and/or differentiation signaling that regulate CSCs, including growth factors (e.g., fibroblast growth factors, TGF- β , bone morphogenetic proteins), cytokines (e.g., interleukins-6 and -8), or morphogens (e.g., Hedgehog, Notch).

These in turn influence many transcription factors including Nanog, Oct4, and SOX2 and c-MYC, which play critical roles in deciding the fate of CSCs [161–163]. In sum, the identification of only four CSC targeting molecules so far using CSC-pathway screening approach for breast CSCs highlights the difficulty of targeting CSCs. For the anti-CSC paradigm to translate into therapeutic success, a more fundamental and generalizable approach is needed.

Concluding Remarks and Opportunities

Our review has identified key research directions in the field of CRC, as summarized below. a) Current genomic testing includes primarily somatic mutations only, and not genetic testing for germline mutations. Adding the latter might allow for more accurate CRC risk assessment, surveillance and prevention. Also, marker(s) such as specific mucins could serve as a powerful adjunct to colonoscopy by enabling efficient detection of SSA/Ps. b) Several emerging genetic biomarkers as well as cell-free miRNAs and small non-coding RNAs have been suggested as noninvasive and highly sensitive candidate markers for CRCs. They will require large clinical trial-based validation. c) In relation to precision treatment approaches, developments in ex vivo cell-genomics technologies and the potential to combine genomic/genetic testing with tumoroid-based drug response testing might provide more accurate clinical efficacy of certain drugs in individual patients. d) Integrating targeted tumor imaging preoperatively and delineating tumor margins intraoperatively with fluorescent labeling can improve the precision of surgical resection. e) Given the pivotal role of cancer-initiating stem cells in CRC progression, the stem cell niches discovered thus far, such as CEACAM/TGF- β and M3R, need further investigation. Other stem cell niches, such as telomerase, CTCF, CEA, E-cadherin, and β -catenin, may be equally important in developing targeted, preventive therapies against CRC. f) CRC tumoroid/organoid technology using polymeric nanofiber scaffolds to expand CSCs ex vivo is of considerable interest. Specifically, tumoroid technology can be used to expand CSCs of hard-to-detect SSA/P neoplasms, which can be characterized and investigated for CRC stem cell niches and used for evaluating existing chemo- and immunotherapies.

Acknowledgments This VA Colorectal Cancer Cell-genomics Consortium (VA4C) meeting was supported by a Department of Veterans Affairs Biomedical Laboratory Research and Development Service field-based meeting award to Dr. Shyam Mohapatra and Dr. Subhra Mohapatra. The contents of any report, written material or manuscripts emanating from this meeting do not represent the views of the Department of Veterans Affairs or the United States Government. The VA4C would like to acknowledge the support of Ms. Christen Bouchard in preparation of this manuscript.

Compliance with ethical standards

Conflict of interest All authors except the following have indicated no financial conflicts of interests in relation to writing of this review. Even though not relevant to this review, Dr. Hemant K Roy has declared that he is a co-founder and shareholder in startup company Nanocytomics and American BioOptics LLC. Additionally, Subhra Mohapatra (co-founder and scientific advisor), Shyam Mohapatra (co-founder and scientific advisor) and the University of South Florida have equity interest in Transgenex Nanobiotech Inc. The terms of the conflict management have been reviewed and approved by the University of South Florida, Tampa, in accordance with its conflict of interest policies.

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
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