Update of the recommendations for the determination of biomarkers in colorectal carcinoma: National Consensus of the Spanish Society of Medical Oncology and the Spanish Society of Pathology

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Abstract
In this update of the consensus of the Spanish Society of Medical Oncology (Sociedad Española de Oncología Médica—SEOM) and the Spanish Society of Pathology (Sociedad Española de Anatomía Patológica—SEAP), advances in the analysis of biomarkers in advanced colorectal cancer (CRC) as well as susceptibility markers of hereditary CRC and molecular biomarkers of localized CRC are reviewed. Recently published information on the essential determination of KRAS, NRAS and BRAF mutations and the convenience of determining the amplification of human epidermal growth factor receptor 2 (HER2), the expression of proteins in the DNA repair pathway and the study of NTRK fusions are also evaluated. From the pathological point of view, the importance of analysing the tumour budding and poorly differentiated clusters, and its prognostic value in CRC is reviewed, as well as the impact of molecular lymph node analysis on lymph node staging in CRC.

Keywords Diagnosis · Neoplastic disease · Gastrointestinal · Genetic prognosis · Tumour

Introduction

During recent years, there have been advances related to the analysis of biomarkers with diagnostic, prognostic and therapeutic value in colorectal cancer (CRC), especially in advanced disease. Five years have passed since the last review of the consensus guide of the Spanish Society of Medical Oncology (Sociedad Española de Oncología Médica—SEOM) and the Spanish Society of Pathology (Sociedad Española de Anatomía Patológica—SEAP) [1], and it is important to update it to incorporate the advances that facilitate the administration of precision therapies to these patients.

In addition, this review examines the novelty of hereditary CRC susceptibility markers and the molecular biomarkers of localized CRC. Recently published information on the essential determination of KRAS, NRAS and BRAF mutations is also evaluated [2], as well as the convenience of determining other biomarkers, such as the amplification of the epidermal growth factor receptor (HER2), the expression of proteins in the DNA repair pathway and the study of NTRK fusions [2].

From a pathological point of view, new data have been published with a high degree of evidence regarding the diagnostic and prognostic value of some biomarkers that will be detailed in this review. One example is the incorporation into pathology diagnostic protocols the analysis of the tumour budding (TB), defined as the presence of isolated tumour cells or small groups of less than five tumour cells at the invasive tumour front, or the poorly differentiated clusters (PDC), defined as groups of five or more tumour cells at the invasive front of the stroma of the CRC which does not form glandular structures. Both, TB and PDC are prognostic factors in CRC [3, 4].

Another important pathological aspect is the introduction of new molecular assays for the analysis of lymph nodes extracted from surgical specimens of CRC. The...
pooling-OSNA® (one-step nucleic acid amplification) method, based on the RT-LAMP (reverse transcription loop-mediated isothermal amplification) of cytokeratin 19 (CK19) mRNA for the detection of lymph node metastases, surpassing the current performance of conventional morphological analysis with H&E. Molecular lymph node staging is relevant, especially in the early stages of the disease [5].

Finally, this review evaluates the future incorporation of new pan-genomic technologies, such as next-generation sequencing (NGS) and liquid biopsy for the clinical management of patients with CRC. All these aspects are developed in this guide, which, like the previous one, will remain open to any necessary revision in the future.

**Clinical aspects**

**Molecular markers of hereditary CRC susceptibility**

Of all CRC cases diagnosed each year, 25% present some characteristic indicating hereditary cancer susceptibility. Of these, 5% correspond to Lynch syndrome, 1% to polyposis and the rest to what could be called ‘family aggregations’.

The identification of these clinical pictures has a high impact on the prevention of CRC, given that the identification of incipient neoplastic lesions leads to cure rates of up to 90% and affects comorbidities, quality of life and health efficiency.

**Lynch syndrome**

The identification of this type of syndrome is based on the Amsterdam I and II criteria [6, 7], which are exclusively clinical. When a family complies, genetic testing should be offered to identify the presence of germline mutations in DNA repair genes (mismatch repair [MMR] genes), such as \(\text{MLH1}, \text{MSH2}, \text{MSH6}\) and \(\text{PMS2}\). This genetic study should always be accompanied by adequate genetic counselling.

The algorithm for the identification of these mutations is well described and accepted by different scientific societies (Fig. 1). Initial screening is performed by identifying the lack of expression of the corresponding proteins of the genes involved in CRC and/or endometrium or by analysing microsatellite instability [MSI]. Any of these techniques is fully accepted today, and each has its specificities. In the case that any of these alterations exist, and once it has been

![Fig. 1 Performance algorithm to detect Lynch syndrome.](image)

- **Assess at the individual level.**
- **When there is no available tumour, but the Amsterdam criteria are met and the implications for the family are important, the option of conducting a study of germline mutations in MMR genes should be evaluated.**
- **When there are no living or available affects, the germinal study should be evaluated in a healthy subject.**

\(\text{MSI}\) immunohistochemistry, \(\text{MMR}\) mismatch repair genes, \(\text{MSI}\) microsatellite instability
ruled out that it is due to an exclusively somatic alteration, the germinal study of the MMR genes should be performed.

Notably, at present, this type of study is being performed in any patient with CRC. This measure is justified not only by the preventive implications for the patient and their families but also because it helps to decide whether to administer adjuvant therapy in patients in whom the indication of such treatment is doubtful [8].

**Polyposis syndromes**

The main characteristic of polyposis syndromes, which usually affect the younger population, is the development of a large number of polyps along the digestive tract, with different histologies and associated with other systemic lesions, with no histological markers to identify them.

If a person and/or their family have a polyposis syndrome, genetic counselling should be offered, and a germinal study of MMR genes should be carried out according to the clinical picture.

Taking into account this information, this panel of experts recommends the following:

- Perform MSI and/or immunohistochemistry (IHC) analysis for all patients diagnosed with CRC; and
- Continue with the germinal study of MMR genes in patients in which MSI is observed and/or loss of expression of repair proteins, provided that an exclusively somatic origin has been ruled out.

**Molecular markers of localized CRC**

The use of adjuvant chemotherapy in patients with stage II CRC is controversial and is only recommended if at least one validated risk factor is observed. Among the clinical, pathological and molecular markers that have been correlated with progression-free survival (PFS) and overall survival (OS), only the evaluation of MSI versus stable microsatellites (MSs), T3 versus T4 and the number of lymph nodes analysed (≥ 12–14 versus < 12–14) have been validated repeatedly in prospective randomized clinical trials [9].

However, in patients with stage III CRC, the prognostic value of MSI is attenuated and is not observed in several large prospective cohort studies. The determination of alterations in RAS and BRAF has been shown to have prognostic value in recent randomized cohort studies but currently has no clinical utility [10].

Several multigenic tests have been developed with a commercial vision, but their overlap is low, and their clinical application is not guaranteed [11]. These tests have focused on patients with stage II CRC, in whom the evidence for administering adjuvant chemotherapy is conflictive, with the aim of selecting patients with higher risk and, therefore, who may benefit more from chemotherapy. Only two tests, Oncotype DX® and GeneFx® Colon [12], have been validated in cohorts of prospective and randomized patients by multivariate analysis in formalin-fixed paraffin-embedded (FFPE) tumour samples. Oncotype DX® is the genetic test with the most evidence and has been developed as a real-time polymerase chain reaction (RT-PCR) multigenic quantitative test. Its prognostic value has been observed in four independent prospective clinical trials that randomized patients with stage II CRC, observing hazard ratios (HR) between 1.43 and 1.68 [13–16]. It is important to note that Oncotype DX® maintains its prognostic value when other clinically relevant prognostic factors, such as T4 or MSI, are included in the multivariate analysis. However, its clinical utility is not guaranteed due to the lack of predictive value of the benefit of chemotherapy, as well as the small, but significant, prognostic differentiation between low, intermediate and high risk.

In this context, Immunoscore® has recently emerged as an alternative prognostic biomarker. It has been developed as a quantitative measure of the presence of CD3+/CD8+ lymphocytes that have invaded a tumour. Both its internal validity and its external clinical validation have been demonstrated in long series of patients with stages II and III CRC through multivariate analysis, including when other factors, such as clinical stage and MSI status, were included in the multivariate analysis.[17]. However, the predictive value of the benefit of chemotherapy is uncertain and lacks independent evidence that demonstrates its prognostic value in stage II CRC, which is only observed when combined with known clinicopathological markers and MSI status.

More promising is the analysis of circulating tumour DNA (ctDNA), through which minimal residual disease can be detected with high prognostic value. If this is validated in the studies of clinical utility that are being carried out, these biomarkers may be the most robust in the near future [18]. Lastly, lack of CDX2 expression confers poor prognosis in stage 2 and may also predict benefit from adjuvant chemotherapy, but this must be confirmed in larger prospective randomized series [19].

**Essential molecular markers of advanced CRC**

CRC is a disease characterized by its high heterogeneity, a product of genomic instability and its interaction with multiple exogenous factors that influence carcinogenesis (diet, lifestyle, microbiome, etc.) and that constitutes the exposome. This genomic heterogeneity translates into different models of carcinogenesis and, ultimately, different phenotypes. Some of these genomic alterations can directly influence the selection of treatments; therefore, it is standard practice to determine the mutational state of multiple genes, including the determination of the extended RAS mutation,
the *BRAF* V600E mutation and the MSI status, which is usually performed by IHC staining for *MLH1, MSH2, MSH6* and *PMS2*.

**RAS mutations**

The main biomarker of clinical utility in metastatic CRC is the mutational state of genes in the *RAS* family (*KRAS/NRAS/HRAS*). Mutations are present in approximately 50% of cases, the most frequent being those that occur in codons 12 and 13 of exon 2 and that condition a state of cellular expansion.

*RAS* mutations are a predictor of the absence of response to treatment with epidermal growth factor receptor (EGFR) inhibitors, such as cetuximab and panitumumab. In the CRISTAL study, the addition of cetuximab to the regimen with folinic acid, 5-fluorouracil and irinotecan (FOLFIRI) showed an increase in the response rate (57% versus 40%; *p* < 0.001) and median OS (23.5 versus 20.0 months; HR 0.796; *p* = 0.0093) in patients with tumours without *KRAS* mutations [20]. However, in patients with tumours with *KRAS* mutations, a 4% decrease in the response rate was evidenced, and there were no significant differences in terms of OS [20]. Similar findings were found when analysing the OPUS trial, in which the chemotherapy regimen was folinic acid, 5-fluorouracil and oxaliplatin (FOLFOX) with cetuximab, and in the PRIME trial, in which FOLFOX was combined with panitumumab [21, 22]. Taking into account these data [23], guidelines from the American Society of Clinical Oncology (ASCO), the European Society for Medical Oncology (ESMO) and the National Comprehensive Cancer Network (NCCN) recommend the extended determination of *RAS* mutations before the administration of EGFR inhibitors [8, 24, 25]. The *RAS* analysis should include *HRAS, KRAS* and *NRAS*, exons 2 (codons 12 and 13), 3 (codons 59 and 61) and 4 (codons 117 and 146) [8, 24, 25].

**BRAF mutations**

The most frequent *BRAF* mutation is the substitution of glutamic acid for valine in codon V600E, which produces constitutive activation of the MAPK pathway. This mutation occurs in 8–10% of patients with metastatic CRC and excludes the *RAS* mutation [26]. It is more common in women, usually appears in more advanced stages and has been associated with tumours of the right colon, poorly differentiated, with mucinous histology and with MSI. Additionally, it seems that *BRAF* mutations lead to the development of more peritoneal metastases and less disease limited to the liver or lung.

Regarding the prognostic role of *BRAF*, it seems that mutation is a negative prognostic factor, although its association with MSI has to be taken into account. A joint analysis of four phase III clinical trials of first-line treatment in patients with metastatic CRC evaluated the possible prognostic value of *BRAF* [27]. In a total of 3063 patients, of whom 8.2% had a *BRAF* mutation, the OS was 11.4 months, and the PFS was 6.2 months, compared to 17.2 months and 7.7 months, respectively, that were observed in patients without *BRAF* mutations (*p* < 0.001). This same study analysed the possible prognostic value of MSI, and it was observed that the OS for patients with MSI of high instability (MSI-H) was 13.6 months, while in patients with MSs, it was 16.8 months (*p* = 0.001). When the *BRAF* and MSI statuses were analysed, there were no differences in terms of PFS or OS in the MSI-H population regardless of *BRAF* status, but in the MSs population with mutated *BRAF*, a significant decrease in OS and PFS was observed. These data support the negative prognostic value of *BRAF* mutations.

Recent data support the predictive value of *BRAF* mutations [28]. Phase III BEACON Colorectal Cancer Study enrolled 665 patients with *BRAF* V600E across three cohorts, including the triplet regimen of encorafenib, biningtinib and cetuximab; the doublet regimen of encorafenib and cetuximab; and a control cohort where patients could receive either cetuximab and irinotecan or cetuximab with FOLFOX. At the time of the interim analysis, the median duration of follow-up was 7.8 months. The median OS was 5.4 months, 8.4 months and 9.0 months for the control group, the doublet regimen arm, and the triplet regimen arm, respectively. Comparing the results obtained in the triplet regimen arm with the control group, HR was 0.52 (*p* < 0.001). An independent central review committee assessed the response rate in this analysis for the first 331 randomized patients. The response rate in the control arm was 2% compared with 20% in the doublet regimen arm and 26% in the triplet regimen arm (control arm versus triplet regimen arm; *p* < 0.001).

**MSI**

Mutations in the *MLH1, MSH2, MSH6* and *PMS2* genes are related to poor *MMR* as well as MSI. They appear in 15% of stage II and III tumours but in only 4% of stage IV tumours. In stages II–III, MSI is related to a better prognosis, and specifically in stage II, the better prognosis occurs in the absence of benefits from adjuvant treatment with fluoropyrimidines. However, MSI in stage IV is associated with an unfavourable prognosis, most likely due to its association with *BRAF* mutations.

Two recent studies show that the alteration of *MMR* may have a predictive role in response to immunotherapy; therefore, the NCCN recommends its determination in all patients with metastatic CRC [29, 30]. The Food and Drug Administration (FDA) has approved the use of nivolumab
and pembrolizumab in patients with MMR-deficient and MSI-H metastatic CRC.

**Recommended molecular markers of advanced CRC**

HER2 is overexpressed in a small proportion of CRC. The Cancer Genome Atlas (TCGA) project identified somatic amplifications or mutations of HER2 in 7% of CRC cases and in 5% of the non-hypermutated CRC subgroup (4% amplification, 3% mutation and 1% both), being more frequent in native RAS tumours (8%) [31]. An analysis of 3256 patients with CRC from different clinical trials (QUASAR, FOCUS and PICCOLO) confirmed a higher incidence of HER2-positive CRC in native KRAS/BRAF tumours and advanced stages [32]. Some studies indicate that the amplification of HER2 is more frequent in distal tumours (PETACC-3 and HERACLES-A), but others have not confirmed that observation (PETACC-8) [33]. Its association with survival is also controversial [32, 33].

Protein overexpression and gene amplification are easily detectable by IHC or fluorescent in situ hybridization (FISH), respectively. The criteria developed and validated in the HERACLES programme defined HER2-positive tumours as tumours with an IHC staining intensity of 3+ in more than 50% of cells or a staining intensity of 2+ with a HER2:CEP ratio (chromosome enumeration probe) greater than 2 per FISH in more than 50% of the cells [34]. Amplification of HER2 is involved in innate and acquired resistance to anti-EGFR drugs [33]. The dual inhibition of HER2 (lapatinib and trastuzumab or pertuzumab), but not the simple block, is capable of inducing durable responses in patient-derived xenograft (PDX) models of native RAS/BRAF/PI3K cetuximab-resistant CRC [33]. Consistent with this, the HERACLES-A trial documented a response rate of 30% in 27 patients with native KRAS CRC, positive HER2, and anti-EGFR resistance treated with lapatinib and trastuzumab [35]. Similarly, in the MyPathway Phase II trial, a response rate of 38% was observed in 37 patients with HER2-positive metastatic CRC treated with trastuzumab and pertuzumab [36]. Other strategies being explored are anti-HER2 monoclonal antibody–drug conjugates, such as trastuzumab-emtansine (HERACLES-B and RESCUE assays), and trastuzumab-deruxtecan, at a 25% response rate documented in a phase I study in 12 patients with HER2-positive CRC [37].

The determination of HER2 may, therefore, be of clinical utility and could be justified at least in native RAS tumours resistant to anti-EGFR treatment.

The constitutive activation of different kinases as a consequence of gene translocations plays an essential role in the tumourigenesis of multiple neoplasias, including a subgroup of neoplasias of the colon and rectum, and their pharmacological inhibition constitutes one of the greatest therapeutic successes in the field of precision oncology. In a recent series, in which 18,407 tumour samples and 513 ctDNA samples from CRC patients were characterized by massive sequencing, genetic rearrangements of potentially treatable kinases were identified in 126 tumours (0.68%) and seven ctDNA samples (1.36%) [38]. The most frequently identified fusions in this series involved the RET (0.15%), BRAF (0.12%), NTRK1 (0.14%) and ALK (0.09%) genes. Ninety percent of the tumours with these gene rearrangements were native KRAS, with a high proportion of MSI-H: 86.4%, 45.5% and 14.3% of the tumours with NTRK1, RET and ALK fusions, respectively. The presence of MSI or the absence of RAS mutations may, therefore, help to better select the subgroup of patients in which it is more profitable to proceed to the molecular screening of these gene alterations.

These fusions can be identified with different methods. Classically, the detection of ALK or ROS1 has been performed using FISH or RT-PCR, but there is a growing trend in the use of mass DNA sequencing techniques, which has the advantage of allowing the simultaneous evaluation of multiple genes with potential utility as therapeutic targets. Other useful techniques include complete or directed sequencing of tumour RNA, sequencing of circulating DNA or IHC (generally used as screening).

The presence of gene rearrangements in ALK, ROS1 or NTRK has been associated with worse survival (15.6 versus 33.7 months for patients with [n = 27] or without [n = 319] tumour gene fusions, respectively) [39]. RET fusions have also been associated with worse survival in a series of 24 positive RET patients [40]. In November 2018, the FDA approved the specific TRK inhibitor larotrectinib for the treatment of tumours with NTRK1/2/3 fusions, with an overall response rate of 75% (including two of three patients with CRC) and a median duration of response and PFS not attained after a median follow-up of 9.4 months (PFS at 1 year: 55%) [39]. This is the second tumour-agnostic approval in the history of antineoplastic therapy after approval of pembrolizumab for MSI tumours. Entrectinib was approved in August 2019 for the treatment of tumours with NTRK fusions.

**Pathological aspects**

**What is relevant in the pathology diagnosis of CRC?**

**Tumour budding (TB) and poorly differentiated clusters (PDC)**

The most important prognostic factors in CRC according to the American Joint Committee on Cancer (AJCC) and the 8th edition of the International Union Against Cancer (UICC) are (i) TNM stage, (ii) venous, lymphatic or
perineural invasion and (iii) discontinuous tumour deposits [41]. The determination of TB has been incorporated into the pathology diagnosis protocol of the College of American Pathologists (CRC) v4.0.0.1 and the NCCN guidelines v2.2019. Other factors, such as type, histological grade, the configuration of the tumour border and intratumoural inflammatory infiltrate, influence the prognosis and treatment of patients. Tumour budding or TB, defined by the presence of isolated tumour cells and/or groups of less than five cells in the stroma of an invasive front of the tumour, is the morphological evidence of the epithelial-mesenchymal transition process and is an independent factor of poor prognosis in CRC. TB is relevant in different clinical contexts: (i) in pT1 CRC as an independent predictor of lymph node metastasis, (ii) in stage II CRC predicts recurrence risk and contributes to therapeutic management [42, 43], and (iii) intratumoural TB in endoscopic biopsies allows the individualization of neoadjuvant treatment in rectal carcinomas [44, 45]. The International Tumour Budding Consensus Conference (ITBCC) agreed to count TB with HE at the area of the pathological tumour diagnosis protocol of the College of American Pathologists (CRC) v4.0.0.1 and the NCCN guidelines v2.2019. Other factors, such as type, histological grade, the configuration of the tumour border and intratumoural inflammatory infiltrate, influence the prognosis and treatment of patients. Tumour budding or TB, defined by the presence of isolated tumour cells and/or groups of less than five cells in the stroma of an invasive front of the tumour, is the morphological evidence of the epithelial-mesenchymal transition process and is an independent factor of poor prognosis in CRC. TB is relevant in different clinical contexts: (i) in pT1 CRC as an independent predictor of lymph node metastasis, (ii) in stage II CRC predicts recurrence risk and contributes to therapeutic management [42, 43], and (iii) intratumoural TB in endoscopic biopsies allows the individualization of neoadjuvant treatment in rectal carcinomas [44, 45]. The International Tumour Budding Consensus Conference (ITBCC) agreed to count TB with HE at the area of the tumour front with the most density using a 20× objective and a field of 0.785 mm². TB is classified as Bd1-low (0–4 buds) with marked acute inflammation. TB must be distinguished from PDCs, defined as groups of five or more tumour cells in the invasive tumour front that do not form glandular structures [4]. TB should not be reported in rectal tumours treated with neoadjuvancy, or in mucinous carcinomas with signet ring cells within mucus lakes. If TB cannot be assessed, it is indicated as no non-assessable with an explanatory note [47, 48].

**Lymph node staging**

Lymph node staging (pN) is an important prognostic factor in CRC, a predictor of relapse and survival and determinant of the therapeutic management of patients. The pN stage is obtained from the analysis of lymph nodes stained with HE, which has low sensitivity for detecting lymph node micrometastases in stages I-II CRC. HE staining analyses less than 1% of lymph node tissue [49–53], resulting in 11–24% false negatives. A meta-analysis concluded that the presence of micrometastases in lymph nodes undetected by HE is associated with worse survival [54]. To improve the sensitivity of lymph node analysis, molecular lymph node staging has been performed, which has shown an improvement in pN staging, with respect to HE.

The OSNA technique detects copies of CK19 messenger RNA (mRNA) in lymph nodes, achieving overstaging in 11–50% of HE-determined pN0 patients. Likewise, the total tumour load (TTL) or number of CK19 copies in the lymph nodes is correlated with classical high-risk factors in CRC, such as pT and pN stage, histologic grade, mucinous or signet ring histology, tumour size, male sex, lymphatic invasion and number of lymph nodes assessed [55, 56]. Recently, TTL has been correlated with prognosis. The incorporation of molecular lymph node staging in CRC will allow to select stage II patients for adjuvant therapy and assist patient management.

**What are the requirements of an optimal sample?**

For the study of biomarkers in CRC, a sample with enough amount of well-preserved tumour will guarantee the performance of the analysis and quality of the results.

**Preservation of tumour material**

Currently, the primary source for biomarkers studies are FFPE tumours and normal tissues, stored at room temperature.

Although the use of formaldehyde solution is the universal method of tissue fixation, formaldehyde produces covalent bridges in aqueous solutions and interacts with the -NH2 groups of proteins, forming cross-linked methylene bridges; the tertiary and quaternary conformations of these bridges are reversible (epitope retrieval), and they interact with DNA by forming hydroxy methylene bridges between two amino groups. In addition, treatment with formaldehyde can produce aberrant mutations by the formation of apurinic and apyrimidinic sites, DNA degradation and cross-linking of cytosines and has recently been considered carcinogenic. Therefore, alternative fixatives have been developed to improve the preservation of nucleic acids. Among these, PAXgene® Tissue Fix (PreAnalytix GmbH Hombrechtikon, Switzerland) and glyoxal acid-free (GAF) stand out [57, 58]. The prolonged storage of paraffin blocks at room temperature causes the constant degradation of the nucleic acids regardless of the type of fixative used. To slow this degradation process, the blocks should be stored at 4 °C or frozen [59].

**Minimum sufficient quantity**

It is crucial to select tumour material through macro- or microdissection to avoid contamination with normal tissue. To guarantee results, a tumour cell content 10% higher than normal cell content is considered optimal, thus avoiding the possibility of false results. Currently, with the latest generation of sequencing technologies, a threshold of 5% could be established.

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The ideal study sample contains the greatest possible number of tumour cells. Thus, it is preferable to compare a resected piece of CRC with the content of a diagnostic endoscopic biopsy or cytological material [60].

Which techniques are most appropriate for the determination of each biomarker?

Table 1 summarizes the most important biomarkers in CRC along with the recommended techniques to evaluate them and whether its performance is optional or mandatory.

RAS mutations

Currently available technologies for mutational analysis of the KRAS and NRAS genes have not been significantly modified compared to the last guide [61, 62], except for the incorporation of the mutational status of the BRAF gene [25], as reflected in Table 2.

A large number of laboratories continue to use molecular techniques based on sequencing as confirmation techniques of mutations (Sanger sequencing and pyrosequencing) or as screening techniques based on NGS, where different platforms allow these determinations to be performed simultaneously in both clinical trials and in research studies as well as in the assistance activities of several molecular pathology laboratories in Europe [63]. However, the vast majority of laboratories have focused their activity on the use of specific assays to detect specific alterations in these genes (KRAS, NRAS and BRAF) using commercial kits by Conformité Européenne-In Vitro Diagnostic (CE-IVD) at the expense of the tests developed by each laboratory (laboratory developed test [LDTs]), whose use is limited in Spain in comparison with other European countries, such as Holland and Germany [64].

Notably, many of the kits used to determine KRAS/NRAS mutations do not determine BRAF mutations simultaneously; therefore, many laboratories have had to modify their workflows as well as combine molecular techniques to analyse these mutations sequentially (e.g., first, KRAS gene mutations are analysed, then NRAS gene mutations and finally BRAF gene mutations), which increases response times, sample manipulation and complexity for obtaining these results.

In the case of the BRAF gene, the majority of the mutations focus on the valine amino acid at position 600, which can be affected in multiple ways (V600E/E2/K/D/R or M); however, given that the vast majority correspond to the p. V600E change, this mutation can be detected by a specific antibody (Clone VE1) via IHC. However, although this technique has shown high sensitivity and analytical specificity in different studies [65, 66], confirmation is currently required by a molecular technique.

In recent years, there has been an increase in the use of commercial point of care (POC) techniques that minimize and automate the process of determining gene mutations, significantly reducing response times, reducing the risk of obtaining false positives/negatives and avoiding DNA extraction because analyses can be performed directly with an FFPE sample, with minimum manipulation by the technical staff and with validated results against conventional

| Biomarker | Technique | Indication |
|-----------|-----------|------------|
| CRC—universal |  |  |
| MSI | IHC and/or MSI analysis by qPCR, bPCR, NGS | Required |
| Immunoscore or immunodensity | IHC and/or digital score | Optional |
| CRC—localized |  |  |
| MSI | IHC and/or PCR and MSI analysis | Required |
| Immunoscore or immunodensity | IHC and/or digital score | Optional |
| CRC—advanced |  |  |
| MSI | IHC and/or PCR and MSI analysis | Required |
| Extended RAS | KRAS, NRAS | Required |
| B-RAF | V600E | Required |
| | V600E2/K/D/R or M | Optional |
| HER2 | IHC / FISH / SISH | Optional |
| Rearrangements of: NTRK1; NTRK2; NTRK3 | IHC and FISH, NGS, RT-PCR, NanoString® | Optional in MSI, MLH1 hypermethylation and RAS WT |
| Liquid biopsy | qPCR, dPCR, NGS, cfDNA Idylla | Optional for patient monitorization |

cfDNA cell free DNA, FISH fluorescent in situ hybridization, IHC immunohistochemistry, MSI microsatellite instability, NGS next-generation sequencing, PCR polymerase chain reaction; bPCR bridging PCR, dPCR digital PCR, qPCR quantitative PCR, RT-PCR real-time PCR, SISH silver in situ hybridization, WT wild type
### Table 2  Molecular techniques employed in CRC

| Techniques available                          | Sensitivity (% mutated DNA) | Characteristics                                                                 |
|---------------------------------------------|----------------------------|--------------------------------------------------------------------------------|
| **Mutational study of isolated genes (KRAS, NRAS and BRAF)** |
| Methods of direct sequencing                |                            |                                                                                |
| Sanger method                               | 25                         | Detects any mutation, Requires a greater amount of mutated DNA, Inexpensive      |
| Pyrosequencing                              | 5–10                       | Commercial trial available, Requires pyrosequencer                             |
| NGS                                         | 1–5                        | Commercial trial available, Requires specific equipment, Requires experience in molecular biology |
| PCR-fragment analysis                       | 1–5                        | Commercial trial available, Requires specific equipment, Requires experience in molecular biology |
| Quantitative RT-PCR                         |                            |                                                                                |
| TaqMan ® PCR                                | 10                         | Only detects specific mutations, No commercial trial, Requires real-time thermocycler |
| Scorpions-ARMS                              | 1                          | Only detects specific mutations, Commercial trial available                     |
| Mutated allele enrichment techniques        |                            |                                                                                |
| PNA-LNA PCR clamp                           | 0,1–1                      | Only detects specific mutations, Requires non-commercial LNA probes, Requires experience in molecular biology |
| COLD-PCR                                    | 0,1–1                      | Requires experience in molecular biology, Can be associated with sequencing and pyrosequencing techniques |
| PCR–RFLP                                    | 5                          | Only detects mutations that generate a restriction site, Commercial trial available |
| DHPLC                                       | 1                          | Detects any mutation, Special equipment required, Requires experience in HPLC    |
| HRM                                         | 1                          | Detects any mutation, Requires specific equipment, Requires experience in molecular biology |
| PCR + hybridization                         | 1–5                        | Only detects specific mutations, Commercial trial available, Requires specific equipment (in the case of hybridization in arrays or strips if there is a large volume), Requires experience in molecular biology |
| Direct techniques for determining RAS       |                            |                                                                                |
| Point of care                               | 1–5                        | Only detects specific mutations, Requires specific equipment, Does not require experience in molecular biology |
| IHQ BRAF (VE1)                              |                            | Only detects the mutation V600E of BRAF, Requires specific automation equipment, Does not require experience in molecular biology |
| **MSI study**                               |                            |                                                                                |
| IHQ                                         |                            | Detects mutations in the MLH1, MHS2, MSH3 and PMS2 genes, Requires specific automation equipment, Has internal controls for each staining |
| PCR + MSI analysis                          | 1–5                        | Detects MSI alterations, Commercial trial available, Requires sequencing and experience in molecular biology |

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techniques and NGS [67], even using DNA extracted from paraffin-embedded samples [68].

In the coming years, it is believed that the use of specific algorithms based on the use of artificial intelligence will increase and be used to analyse both radiological and HE images, which will produce RAS analyses with greater precision [69, 70].

**MSI**

Other indispensable markers in patients with CRC are MMR deficiency and the presence of MSI for identifying patients with Lynch syndrome with prognostic purpose and to predict the response to different immunotherapeutic treatments.

For this purpose, in the majority of pathology laboratories, IHC analysis of the four repair genes (*MLH1, MSH2, MSH6* and *PMS2*) is performed; however, because the encoded proteins act in tandem, different groups have proposed that only IHQ determination of *MLH1* and *MSH2* can be used as a screening technique [71].

In addition, PCR techniques have been used for the detection of MSI. When developing the methodology, we began using a panel of microsatellites (Bethesda panel) composed of markers that detect single base alterations (BAT25 and BAT26) as well as three dinucleotide markers (D2S123, D5S346 and D17S250), but recently, new mononucleotide satellites have been incorporated, which have shown better prediction capabilities than those of previous dinucleotides [72].

Finally, both recently implemented NGS techniques and new POC tools allow the determination of MSI with new satellites included in their panels and are being validated in different studies [73].

**Alterations in HER2**

The determination of the amplification of HER2 in CRC has generated great interest because of its prognostic and predictive value of response to specific treatments because approximately 3% of patients have such alterations [74].

The methods of analysis are the same as those used in breast and gastric cancers, where IHC (HercepTest® or Clone 4B5) stands out, showing great concordance with the gene amplification detected by FISH or in silver in situ hybridization (SISH) [34]. Currently, both the determination of the copy number alterations (CNAs) in a tumour by

| Techniques available | Sensitivity (% mutated DNA) | Characteristics |
|----------------------|-----------------------------|-----------------|
| NGS                  | 1–5                         | Available commercial trials pending validation Requires specific equipment Requires experience in molecular biology |
| **HER2 amplification study** |                          | High concordance with gene amplification in those overexpressed Requires specific automation equipment Has interpretation guides |
| IHQ                  | 1–5                         | Detects amplifications in a specific way Commercial trials available Requires experience and fluorescence microscope (FISH) |
| In situ hybridization | 1–5                         | Detects amplifications in a specific way Commercial trials available Requires experience and fluorescence microscope (FISH) and the use of three probes |
| NGS                  | 1–5                         | Available commercial trials pending validation Requires specific equipment Requires experience in molecular biology |
| **NTRK rearrangements study** |                          | Overexpression associated with the presence of rearrangements of any of the three genes (*NTRK1, NTRK2* and *NRTK3*) Requires specific automation equipment Confirmation with additional molecular techniques |
| IHQ                  | 1–5                         | Detects amplifications in a specific way Commercial trials available Requires experience and fluorescence microscope (FISH) and the use of three probes |
| In situ hybridization | 1–5                         | Detects amplifications in a specific way Commercial trials available Requires experience and fluorescence microscope (FISH) and the use of three probes |
| NGS                  | 1–5                         | Available commercial trials pending validation Requires specific equipment Requires experience in molecular biology |

ARMS mutation system refractory to amplification, COLD-PCR coamplification at lower denaturation temperatures, DHPLC denaturing high-performance liquid chromatography, FISH fluorescent in situ hybridization, HPLC high-performance liquid chromatography, HRM high-resolution fusion, IHC immunohistochemistry, MSI microsatellite instability, NGS next-generation sequencing, PCR polymerase chain reaction, PNA-LNA peptide nucleic acid-blocked nucleic acid, RFLP restriction fragment length polymorphisms, RT-PCR real-time PCR

Table 2 (continued)
NGS techniques and the quantification of the levels of HER2 mRNA are being validated in different studies before their implementation in routine healthcare [39].

**NTRK rearrangements**

The rearrangements of the \(NTRK1, \ NTRK2\) and \(NTRK3\) genes are of special relevance in CRC because, although their prevalence is very low (1.5%) [39], these alterations are associated with the presence of MSI, hypermethylated \(MLH1\) and native \(RAS\) [75]. In addition, it is important to mention that they have a great predictive value of response to TRK inhibitors. The classical techniques for the detection of \(NTRK\) alterations are generally performed by IHC with pan-TRK clones as screening, with subsequent confirmation by different tests (FISH, NGS, RT-PCR or NanoString®); however, each one has its limitations [76].

**What is the current and future role of NGS in CRC?**

The implementation of NGS in the study of CRC will allow, in the future, the diagnosis of Lynch syndrome through the mutational study of the \(MMR\) and \(EPCAM\) genes [77] and will guide treatment by detecting the mutations of the main genes involved in the pathways of CRC carcinogenesis, such as \(RAS\) and \(BRAF\), as well as the genes associated with targeted treatment, such as \(HER2\). In addition, it will be possible to classify CRC into molecular subtypes (CMS1, CMS2, CMS3 and CMS4) with different biological behaviours and therapeutic responses, which will open the possibility of designing personalized treatment [9]. Likewise, NGS will allow the identification of the tumour hypermutation status, which will guarantee the efficacy of immunotherapy [78].

Furthermore, through the study of ctDNA, it will be possible to monitor response to treatment, anticipate the appearance of local recurrence and metastasis, and detect resistance to ongoing treatment [79].

Finally, the possibility of implementing the combination of different molecular tools in the diagnostic routine will also improve the quality of tumour samples [68], without forgetting that IHC continues to be an indispensable diagnostic technique [80].

**What are the current and future roles of liquid biopsy in CRC?**

A liquid biopsy is based on the analysis of a biological liquid, usually blood, but also cerebrospinal fluid or urine, to reveal characteristics of a cancer, such as circulating tumour cells (CTCs), free nucleic acids (DNA or RNA), exosomes or tumour-derived platelets. The quantification of CTCs in peripheral blood has been shown to have prognostic value with a threshold level above or below three cells. The possibility of extracting DNA from these cells for NGS has been recently evaluated [81].

The most developed option in liquid biopsy is the study of ctDNA in peripheral blood. The biomarker with the greatest impact on CRC is the mutation in the \(RAS\) oncogene, including \(KRAS\) and \(NRAS\), in exons 2, 3 and 4, which indicates patients who may be candidates for treatment with anti-EGFR antibodies in case of the native or non-mutated sequence. This technique has been studied using tumour tissue as a comparison. Different publications have confirmed that using high sensitivity molecular techniques [82], the concordance between the analyses performed on tumour tissue and peripheral blood exceeds 90% [83]. Correctly selecting the analysis technique is critical. Liquid biopsy has two very significant advantages compared to tissue biopsy: (i) it indicates the heterogeneity of the tumour and (ii) it is a real-time reflection of the molecular profile, which allows determining the dynamic evolution of the tumour.

Several studies have observed the convenience of using liquid biopsy in the early determination of response in patients with \(RAS\) mutations, in whom during the course of treatment the basal mutation was not detected, and in patients with a \(RAS\) native sequence, in whom during treatment a \(RAS\) mutation is detected, predicting, presumably, the onset of radiological progression of the disease. The determination of \(BRAF\) mutations can be another important application of liquid biopsy because it allows monitoring patients with CRC with this alteration [84].

Recent research has opened the way to new uses of liquid biopsy. Some studies have proposed its possible use for the early diagnosis of CRC; however, this option is still very preliminary and, for the moment, lacks an adequate cost-effectiveness relationship as a population screening technique [85]. Finally, the identification of low-incidence molecular alterations in CRC, such as the presence of \(HER2\) amplification or fusions that affect \(NTRK\), could be evaluated via ctDNA by mass sequencing platforms [86].

**Conclusions**

In recent decades, significant progress has been made in understanding the molecular characteristics of CRC and in the identification of specific mutations that have allowed the development of new prognostic and predictive biomarkers in the different stages of the disease. These advances, together with those achieved through pathological knowledge, have justified a new edition of this guide.

With respect to the susceptibility markers of hereditary CRC, this panel recommends the analysis of MSI and/
or IHC in all patients diagnosed with CRC. In addition, the germinal study of MMR genes should continue in patients with MSI and/or loss of repair protein expression. In localized stages, the determination of MSI as a prognostic marker is required, especially in stage II CRC. The standardized implementation of genomic platforms is not recommended due to the lack of predictive value of response to chemotherapy.

In patients with advanced CRC, it is essential to determine extended RAS mutations, the BRAF V600E mutation and the MSI status, which is usually performed via IHC for MLH1, MSH2, MSH6 and PMS2. The determination of extended RAS mutations is mandatory before making therapeutic decisions (I-A). The RAS analysis should include HRAS, KRAS and NRAS, exons 2 (codons 12 and 13), 3 (codons 59 and 61) and 4, as negative predictive factors of the anti-EGFR response. The BRAF V600E mutation is required due to its recognized negative prognostic value (I-A) and, recently, as a predictive marker of specific biological treatments (cetuximab, encorafenib and binimetinib). The determination of MSI acquires predictive value in advanced CRC for immunotherapy treatment with pembrolizumab or nivolumab (II-B).

Another recommended marker in advanced CRC is the amplification of HER2 as a negative predictive marker of anti-EGFR response and as a response marker for dual anti-HER2 therapy. Its determination is recommended, at least in native RAS tumours resistant to anti-EGFR. The ALK, ROS1 and NTRK fusions, despite their low incidence, have clinical interest due to recent approval for specific treatments.

The greatest contribution to the genetic study of CRC will come from liquid biopsies, with which minimal residual disease can be detected after surgery. This technique reflects tumour heterogeneity and molecular profiles in real time, reporting the dynamic evolution of the tumour. The implementation of NGS will improve genotypic knowledge, the tumour heterogeneity and molecular profiles in real time, disease can be detected after surgery. This technique reflects otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

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Compliance with ethical standards

Conflict of interest The authors declare that, when writing and revising the text, they did not know the names of the pharmaceutical companies that provided financial support for this project, so this support has not influenced the content of this article.

Ethical approval (research involving human participants and/or animals) The study has been performed in accordance with the ethical standards of the Declaration of Helsinki and its later amendments. This article does not contain any studies with human participants or animals performed by any of the authors.

Informed consent For this type of study formal consent is not required.

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