Colorectal cancer (CRC) is one of the most common malignancies worldwide. In recent years novel therapies targeted at EGFR receptor have been developed. However, this treatment can only be beneficial if no mutation in specific loci of KRAS/NRAS and BRAF genes is found in tumour specimen. Therefore, clinically useful pathological diagnosis of CRC in the era of personalised medicine is a multistep procedure, requiring good cooperation between the clinician/surgeon, pathomorphologist, and molecular biologist. Herein we propose the guidelines of colorectal cancer operating material proceedings for clinicians and pathomorphologists, which determines the correct pathomorphological diagnosis, and we discuss the colorectal cancer molecular biology issues useful in the selection of individual molecular targeted therapy. We discuss and stress the importance of each diagnostic phase: from tumour resection and sample collection at preanalytical stage, through proper pathological preparation, evaluation and selection of material for molecular testing, to molecular analysis and finally preparation of a pathological molecular report.

**Key words:** colorectal cancer, KRAS, NRAS, BRAF, diagnostic, mCRC.

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## Quality and practical aspects of pathological and molecular diagnostics in metastatic colorectal cancer (mCRC)

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### Introduction

In 2013, colorectal cancer (CRC, including colon and rectal cancers) was the second most common female cancer (7173 cases, accounting for 9.2%), and a third of all tumours diagnosed among males (8726 cases, 11.2%) in Poland. Furthermore, CRC contributed to the demise of 5851 men and 4711 women, which made it the second and the third cause of cancer-related deaths in 2013, respectively. According to global data, in 2012, colorectal cancer was diagnosed in a total of 1.4 million patients and was the third most common cancer worldwide [1].

The methods of colorectal cancer treatment are determined by several factors, of which tumour location (colon vs. rectum) and staging according to TNM, are the most important. In recent years treatment targeted at the selective inhibition of tumour cell growth using small molecule inhibitors and monoclonal antibodies has been introduced. This method of therapy is largely based on the knowledge of tumour biology. Hence, the challenge for modern molecular diagnostics is to enable precise selection of patients, who can benefit from different treatment strategies.

Targeted therapies in colorectal cancer are primarily directed at blocking the epidermal growth factor receptor (EGFR receptor), which is involved in signalling pathways controlling cell proliferation, angiogenesis, apoptosis, and metastasis. Overexpression of the receptor, observed in 45–80% of patients with CRC, is associated with worse prognosis: a higher cancer grade, lymph node involvement, and a higher frequency of metastasis to the liver and to lungs, bones, etc. [2–4].

In colon cancer cells, especially in the deepest layers of the tumour, besides EGFR overexpression, higher concentration of receptor-activating ligands is also observed. Importantly, during activation of the receptor, signalling crosstalk and cross-phosphorylation may occur. This allows for the mutual activation of neighbouring receptors or interactions between different signalling pathways. During neoplastic transformation, the self-activation processes, in which the cell itself produces ligands for receptors on its own surface, might be possible as well [4–6].

Currently, the most efficient blocking of EGFR-dependent pathways is achieved using two monoclonal antibodies: cetuximab and panitumumab, which bind the extracellular domain of the EGFR receptor, preventing its dimerisation and self-activation. As a result, signalling transduction, synthesis, and secretion of proangiogenic factors are reduced, the cell cycle is arrested, and pro-apoptotic processes are strengthened. These agents also
Pathologists, and molecular biologists. All diagnostic should, in particular, be followed by clinicians/surgeons, using molecular tests, evidence-based recommendations and/or molecular techniques that enable detection of somatic KRAS, NRAS and BRAF genes.

Monoclonal antibody therapies targeting epidermal growth factor receptor (anti-EGFR therapy) are currently used in Poland as a first-line treatment for patients with advanced CRC. Cetuximab and panitumumab may be used in monotherapy or in combination with chemotherapy.

Importantly, anti-EGFR therapy can only be included after confirmation of CRC diagnosis by pathomorphological examination and exclusion of mutations in selected regions of KRAS, NRAS, and BRAF genes.

Mutation status of those genes is carried out using molecular techniques that enable detection of somatic changes in FFPE (formalin-fixed and paraffin-embedded) tumour samples. Because the percentage of cells containing mutations in the examined material can be very low, high-sensitivity detection methods of molecular diagnostics can only be used. Similarly, proper preparation and precise pathomorphological characteristics of the colorectal cancer histopathological samples for molecular diagnostics are critical.

The role of pathomorphologist in contemporary oncological diagnostics is to recognise cancer, determine its histological type, degree of maturity, and stage of advancement (pTNM), and to evaluate prognostic and predictive factors. Thus, pathomorphological diagnosis is crucial with respect to the decision of treatment strategy in an individual patient. Consequently, the full clinically useful pathological diagnosis of CRC in the era of personalised therapy depends on good cooperation between the clinician/surgeon, pathomorphologist, and molecular biologist.

Hence, the aim of this paper is to:
• propose the guidelines of colorectal cancer operating material proceedings for clinicians and pathomorphologists, which determines the correct pathomorphological diagnosis,
• discuss the colorectal cancer molecular biology issues useful in the selection of individual molecular targeted therapy.

The pathological and molecular analysis of the tumour tissue

The European Society of Pathology and Royal College of Pathologists has recently elaborated the guideline for laboratories performing molecular pathology for cancer patients [8]. In the new situation in oncology when personalised medicine requires evaluation of predictor factors using molecular tests, evidence-based recommendations should, in particular, be followed by clinicians/surgeons, pathologists, and molecular biologists. All diagnostic procedures, including DNA and RNA extraction, can and should be standardised and verified on a regular basis.

The diagnostic process of cancer specimens includes:
• the preanalytical stage,
• the diagnostic stage performed by the pathologists,
• the molecular analysis conducted by the molecular biologists,
• the final diagnosis prepared as a pathological molecular report.

The preanalytical stage

In the first step of the pre-analytic stage the tumour samples are taken from patients by the endoscopist or surgeon, are sent to the pathological laboratories, and are preserved and fixed.

The clinician who send the material for pathomorphological and/or molecular examination is obliged to correctly mark the container with the material transferred for examination and to deliver all the necessary information about the patient and the specimen. Complete clinical information is crucial for the pathological diagnosis.

A referral for a pathomorphological examination should contain, at least:
• personal data of the patient: name, surname, date of birth, PESEL or PID number marked with the traditional method or barcode,
• clinical diagnosis,
• anatomical location and type of surgical procedure with the marked surgical margins,
• in selected cases, a request for additional immunohistochemical or molecular tests for prediction factors assessment.

The pathologists receive different tissue material from the metastatic colorectal cancer cases: small biopsy from the primary tumour or from metastasis to the liver in inoperable cases or primary tumour specimens after surgery. The small biopsies are fixed rapidly in neutral buffered formalin (4% formaldehyde) by endoscopists. Concerning the large specimen size after surgery, sending the fresh material to the laboratories is the most convenient. Because the tumour tissue requires controlled fixation, proceeding in the pathology laboratory is the optimal method. If the pathology laboratory is located in the hospital, non-fixed postoperative material should be sent there immediately. If the estimated transport time to the pathology laboratory is more than an hour, a vacuum is a preferable method for preservation of the fresh specimens. When the transport is expected to be longer, the surgeons should preserve and fix the specimen by themselves immediately after surgery in the operating theatre. If surgical material is sent to another remote location, the fixing conditions and time, together with transport procedure, must be observed by clinicians.

The surgeons should be familiar with the methods of preservation and fixation of large intestine tumour specimens. The preservation of the postoperative material includes cutting the large intestine along the long axis and unfastening it on an expanded polystyrene, wooden, paraffin, or cork board. The prepared specimen is fixed in...
10% buffered formalin (4% formaldehyde aqueous solution buffered to pH 7.2). Formalin is a commercially available fixative fluid, which is chemically a 40% solution of formaldehyde in water.

Formalin buffering should be obligatory because the formaldehyde sold commercially is often contaminated with various chemical substances and is acidic. This may adversely affect its fixing properties, hamper diagnostic processes, and result in the degradation of genetic material (DNA, RNA). In the case of large surgical specimens, a major problem for molecular pathology is a cold ischaemia occurring between removal of a tissue and its fixation, which alters levels of gene expression (at the RNA and protein level).

The specimen should be fixed in formalin for 48 hours or, preferably, for 72 hours. Longer and better fixation of the material allows removal of thinner sections, enabling better investigation of the extent of cancer infiltration. Proper fixation of the material should be achieved when 10 fixer volumes per one volume of the tissue material are provided. Formalin penetrates the tissue at a rate of 1 mm/h to a depth of up to 5 mm, thus material cutting ensures proper access to tissue penetration. Control of fixation time and temperature is recommended for molecular analysis. It should be emphasised that autolytic changes of the tumour occur if the uncut large intestine is inserted into the vessel, containing formalin volume less than required, which results in the inability to obtain the correct results of molecular tests. Poor fixation of the postoperative material is the first cause of the erroneous results of molecular analysis. The material destroyed at this stage is unrecoverable and it is impossible to correct the pre-analytic stage errors.

In the next step of the pre-analytic stage, the following procedures are performed:

a) macroscopic examination and collection the representative specimens by a pathomorphologist or doctor during specialisation with the participation of a laboratory technician/diagnostician,
b) placement of collected samples in the cassettes and signing them with the test number (entering bar codes in the LIS is suggested) by a laboratory technician/laboratory diagnostician manually or automatically with a barcode,
c) carrying out processing of the material in a tissue processor (reagents and handling times need to be checked),
d) paraffin embedding, slicing, handwriting, or automatic bar code signing and staining by a laboratory technician/diagnostician; it is necessary to control the appropriate paraffin temperature.

It should be emphasised that the tissue material should be processed through subsequent stages (points b, c, and d) in the pathology laboratory ensuring automation of the whole process. Equipment ensuring control of its conditions determines the appropriate quality of tissue material useful both for microscopic assessment and molecular testing. This is the important and obligatory condition affecting the quality of the obtained results. Hand-processing of tissue material should not be allowed during the preparation of histopathological preparations (Figs. 1 and 2).

The diagnostic stage

The final pathological diagnosis in colorectal carcinoma cases should be useful for clinicians in making decisions about treatment of patients. Molecular report in the era of personalised medicine should be integrated with the histopathological report. Pathologists use the synoptic reporting form based on a structured checklist. The checklist format guarantees the completeness of reporting for surgical pathology cases and production of the standardised clinical documentation. Pathologists can provide complete reports based on the College of American Pathologists (CAP) Cancer Protocols and electronic Cancer Checklists. The evolution from narrative reporting to pathology reporting using a configurable templates structured synoptic report...
reduce the risk of lacking important, clinically useful information. A systemic review showed that synoptic reporting improved the completeness of pathology diagnosis and quality of pathology evaluation for solid tumours [9]. Besides traditional parameters for colorectal diagnosis, such as tumour type, grade, invasion depth (pT), and nodal status (pN), other important clinical features are reported, e.g. resection margin, type of local spread (vascular, lymphovascular, perineural invasion), tumour deposits, and degree of tumour regression after preoperative treatment. Synoptic reports also include information about the mean number of lymph nodes examined. The quality of pathology reports is presented by the number of lymph nodes assessment (minimum 12 lymph nodes) and the proportion of the lymph nodes with carcinomatous metastasis to lymph nodes without metastasis.

Table 1. The factors important to consider in making a decision about treatment according to the Eighth Edition of the TNM AJCC/UICC [8]

<table>
<thead>
<tr>
<th>Factor important to consider in making decisions about treatment</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Serum carcinoembryonic antigen (CEA) levels in patients after surgery</td>
<td>CEA level may be measured in blood, plasma, or serum by an enzyme-linked immunosorbent assay (ELISA). It is recommended for monitoring the treatment. CEA levels may be used as a response marker for treatment of IV stage disease</td>
</tr>
<tr>
<td>Tumour regression score in rectal carcinoma after pre-operative treatment</td>
<td>The pathological response to preoperative radiotherapy (rectal cancer), chemoradiation (rectal cancer) or chemotherapy (colon and rectal cancer) is an important prognostic parameter assessed by the pathologists according to score system</td>
</tr>
<tr>
<td>Circumferential resection margin (CRM)</td>
<td>CRM is measured in millimetres between the deepest margin of tumour tissue and the non-serosalised area of colorectum</td>
</tr>
<tr>
<td>Lymphovascular invasion (LVI)</td>
<td>In colorectal specimens the invasion of small and large vessels is associated with lymph node metastasis. The extramural venous invasion is an independent adverse prognostic parameter and risk factor for liver metastasis</td>
</tr>
<tr>
<td>Perineural invasion (PNI)</td>
<td>The invasion of nerves and perineural spaces by the tumour cells correlate with a poor prognosis</td>
</tr>
<tr>
<td>Microsatellite instability (MSI)</td>
<td>MSI is a prognostic factor and predictive for lack of response to 5-fluorouracil (5-FU) chemotherapy. The MSI is a key to diagnose the hereditary nonpolyposis colorectal carcinoma (HNPCC) or Lynch syndrome. The sporadic colorectal cancer with MSI has epigenetic promoter methylation analysis of the MLH1. BRAF mutation and MSI in sporadic colon carcinoma predicts worse prognosis in stage III and IV colon cancer</td>
</tr>
<tr>
<td>KRAS and NRAS mutation status</td>
<td>The status of both genes is a predictive factor and correlates with lack of response to anti-EGFR therapy</td>
</tr>
<tr>
<td>BRAF mutation</td>
<td>BRAF p.Val600Glu mutation in colorectal carcinoma blocks the effect of anti-EGFR therapy and predicts progression in stage IV colorectal carcinoma</td>
</tr>
</tbody>
</table>

Molecular pathological reports join the histopathological macro- and microscopic parameters with prognostic and predictive biomarkers. The colorectal pathological reports include molecular factors evaluated using immunohistochemical and molecular methods. The mutation status of KRAS, NRAS, and BRAF genes and microsatellite instability (MSI) are important biomarkers in the selection of the patients for individual therapy.

The Eighth Edition of the TNM American Joint Committee on Cancer (AJCC) conformed the staging proposed by the TNM Committee of the International Union Against Cancer (UICC) [10, 11], which transformed the traditional version of the diagnostic reports into a more useful form for individualised therapy purposes. The main idea of the new version of TNM system is the term “prognostic stage groups” analysed separately for the tumour of the different anatomic site [12]. The pathological staging in colorectal cancers is based on the microscopic examination of the resected specimen (pTNM). The parameters defined for category T (tumour), N (lymph nodes status), and M (distant metastasis) are required for stage grouping.

Additional factors recommended for clinical care are evaluated as tumour deposits (TDs). In the Seventh Edition of TNM TDs are classified as category pNlc. TD is defined as a nodule of regular or irregular contour of cancer cells situated in pericolic/perirectal fat tissue or in adjacent mesentery without lymph node metastasis. This microscopic parameter changes the stage of the tumour. This microscopic parameter changes the stage of the tumour and is classified as pNlc category. The factors that are important to consider in making a decision about treatment in colorectal cancer cases constitute another group of prognostic factors according to the Eighth Edition of the TNM AJCC/UICC. These parameters are clinically significant.
when the physician and patient make a decision concerning treatment. Finally, the anatomical site of the tumour, histological type, grade, and pTNM should be considered together with these additional factors. Table 1 presents the factors to consider in making a decision about treatment according to the Eighth Edition of the TNM AJCC/UICC.

An additional part of the pathological report is dedicated to immunohistochemical and molecular analysis:

• immunohistochemical testing for mismatch repair (MMR) proteins (MLH1, MSH2, MSH6, and PMS2), molecular analysis of MMR genes using mononucleotide panel (BAT-25, BAT-26, NR-21, NR-24, and MONO-27), or other and MLH1 promoter methylation analysis,
• KRAS, NRAS, and BRAF mutational analysis.

The molecular analysis

Preparation of material for diagnostic testing

In the case of metastatic colorectal cancer, metastasis tumour (e.g. to the liver or lymph nodes) should be tested preferentially. However, if unavailable, the primary tumour tissue may be used.

In each case one of the most serious problems associated with the determination of somatic mutations in cancer tissue is its heterogeneity. The FFPE tumour samples commonly contain a significant percentage of normal cells, tumour microenvironment cells, such as activated fibroblasts, activated macrophages, lymphocyte and endothelial cells of the blood, and lymph vessels.

This issue is not a major concern in the aspect of histopathological diagnosis, but it limits significantly the possibility of isolating separated cancer cells for molecular examination, which can further lead to an erroneous test result. If the number of normal cells is much higher than the number of tumour cells, the signal from mutant DNA is “dimmed” by wild type DNA from normal cells, leading to a false negative result. In order to avoid this, the analysed sample should contain not less than 50% of neoplastic fixation to obtain a reliable result. Use of samples containing smaller proportion of tumour cells is permitted, although in such cases highly sensitive analytical methods, such as quantitative polymerase chain reaction (qPCR), should be used. However, according to recommendations, material for molecular purposes should contain tumour cells (almost) exclusively [13].

Therefore, a key element in the preparation of the sample for molecular analysis is the verification of the material by a pathologist, who: 1. selects a slide containing the highest possible percentage of cancer cells in relation to the normal ones and 2. marks the area of tumour cells on the slide (Fig. 2).

In order to obtain a high percentage of cancer cells, manual separation of normal and cancerous tissue (macrodisssection) on a slide is an option. The FFPE block can be also molten, so that the entire tumour tissue fragment can be macrodissected and subsequently re-embedded in paraffin. Alternatively, microdissection can be performed using automated microdissectors. Such devices, usually coupled with scanners and microscopes, cut-out previously selected fragments of tissue sample from the slide and transfer it to the test tube. Laser micro-dissectors are even able to extract single cancer cells.

The number of sections should depend on the size of the cancerous tissue. In the case of large fragments (diameter of the section about 1 cm), sections (of thick not exceeding 4-5 μm) from 2-3 slides would be sufficient, while a minimum of five slides of small sections (with a diameter of 2-3 mm) is required.

The thickness of the sections usually ranges from 4 to 10 μm. Thin sections allow faster isolation of genetic material, while thicker ones facilitate manual transfer to test tubes. Importantly, the pathologist is not able to evaluate the cells in the deeper layers of thicker sections and verify the presence of the tumour cells there. Furthermore, all tumour material in a block can be unintentionally used and will no longer be available for any future analyses.

It is also crucial to underline that the HE preparation for the assessment of neoplasm percentage must be made just before the preparators for molecular analysis; otherwise, the tumour content on the HE slide and slide(s) for molecular test may be dramatically different, which may negatively affect molecular test results.

Nucleic acids extraction, quantification, and quality assessment

Nucleic acid extraction from paraffin-embedded tissue should be efficient and enable purification from inhibitors of enzymatic reactions.

Currently, dedicated commercial kits seem to be the best choice. There are several sets available, both manual and compatible with automatic isolators. Extraction systems are usually based on columns containing solid phase binding DNA, or they use the technology of magnetic microspheres that capture DNA molecules and can be easily separated from the solution using magnetic stand.

The quantity and quality of isolated genetic material need to be evaluated prior to the analysis.

For this purpose, fluorimetric systems measuring the amount of DNA-binding substance should be used instead of spectrophotometers. The isolated form FFPE often contain many compounds that may interfere with the spectrophotometric absorbance measurements. This is of particular importance when the analysis is based on qPCR technology or next-generation sequencing, which require very precise amounts of nucleic acids.

Moreover, quantitative and quality assessment of DNA based on quantitative (qPCR) reaction might be another option because it also enables assessment of the degree of DNA degradation and hence may indicate the necessity to resign from further analysis (if DNA is heavily degraded) and to repeat the isolation process from another slide.

Turnaround time in the case of molecular diagnostics depends on the mutation detection method. Real-time PCR-based methods are generally rapid because they demand one-step analysis. Sequencing methods, both classical (Sanger) and especially next generation (NGS), require much more time due to the multistep processes. While choosing the method of analysis, the following aspects should be a matter of careful consideration: availability of equipment...
and technical support, limitations of method optimisation, sensitivity and specificity of the method, time necessary to complete analysis, and unambiguity of the results.

**Final stage**

The complete diagnosis, that includes a pathological report and results of molecular tests for predictors, finishes the diagnostic process of cancer.

**Molecular predictors of response to targeted therapies in metastatic colorectal cancer**

**Proteins involved in the response to therapies targeted at mCRC**

RAS proteins belong to the family of G proteins, which have an ability to bind and hydrolyse guanosine triphosphate (GTP). Binding the ligand by the EGFR receptor leads to the activation of the RAS family proteins, which are active when bound to GTP. Activated RAS conducts the signal further to the effector proteins, which include:

- family of RAF serine-threonine kinases: c-RAF, ARAF, and BRAF, which control cell proliferation and differentiation,
- 3-phosphatidylinositol kinase (phosphoinositide 3-kinase – PI3K), responsible for apoptosis regulation and cell survival,
- RALGDS, RGL, and RGL2 and TIA1M1 proteins that activate RAL and RAC proteins; the RAL protein is responsible for the intracellular vesicular transport, while the RAC protein is involved in the cytoskeleton modelling,
- Ca phospholipase (PLCε), involved in calcium-dependent signal regulation control.

RAS protein is inactivated through interaction with GAP proteins (GTP-ase-activating proteins), e.g. neurofibrin 1 (NF1) or p120. NF1 binds to the active RAS and activates its internal GTPase activity. As a result, the GTP hydrolysis process is a thousand-fold accelerated. Conversion of the RAS-GTP to RAS-GDP (inactive form) results in the inhibition of signal transduction.

The RAS subfamily consists of HRAS, NRAS, KRASA, and KRASB proteins, which are encoded by three distinct genes: HRAS, NRAS, and KRAS (encoding two isoforms: KRASA and KRASB – Fig. 3). Mutations in NRAS and KRAS genes have prognostic significance in the context of CRC therapy. At the protein level, mutation in these genes result in the abolition of the internal GTPase activity, making the protein insensitive to GAPs. Consequently, mutated RAS proteins remain active and their accumulation leads to constant cell stimulation. Therefore, such mutations are referred to as activating [14–20].

The majority of activating mutations in NRAS and KRAS genes occur in codons 12 and 13, which accounts for 70%, and 20% of all mutations identified in colon cancer, respectively. Mutations affecting codons 59, 61, 117, 146, and others are rarer (Table 2). RAS mutations are monoallelic changes occurring at the early stages of carcinogenesis (early and mid-advanced adenomas). However, in around 4% of cases mutations in codon 12 and 13 in tumour cells are identified simultaneously [21–24].

**BRAF** is another protein that has been indicated to play an important role in determining the response to targeted therapy in CRC. Activation of the BRAF protein, following phosphorylation by RAS kinase, leads to activation of another signalling protein – MEK kinase – and subsequently ERK kinase. ERK regulates gene expression and cytoskeleton rearrangement and is involved in several cellular processes: metabolism, proliferation, differentiation, aging, and apoptosis. In the inactive form, the conformation of BRAF protein’s kinase domain prevents the binding of ATP (and the activation of MEK kinase). Structurally, this conformation is preserved by trapping the regulatory part of the enzyme, the activation segment, through hydrophobic interactions with the P loop. Indeed, the majority of activating mutations occur in loci involved in interaction of P loop and activation segment, which is critical to maintain the inactive form. As a result of mutation, the inactive form loses its stability in favour of the activated form [28, 29].

Mutations in **BRAF** occur in 5–22% of colorectal carcinomas. More than 40 different **BRAF** mutations in 24 different codons have been identified so far. In CRC, the most frequent (90%) of them is the substitution of valine at position 600 with glutamate, which localises in the kinase domain. In in vitro kinase activity studies, the p.Val600Glu mutant is over 500 times more active than the normal form [28, 29].

Mutations in **BRAF** occur in the early stages of neoplastic transformation and are not capable of initiating the neoplastic process per se. However, for patients with colorectal cancer treated with cetuximab and panitumumab, the p.Val600Glu mutation is a negative predictor. Patients who have normal RAS genes and the mutant form of **BRAF** do not respond to treatment with anti-EGFR antibodies, and have shortened PFS and OS as compared to patients with wild type **BRAF** [30].

![Fig. 3. The KRAS gene is located on the shorter arm of chromosome 12 (12p1 locus). It consists of six exons. The first exon, (grey) is not translated. Coding exons (blue) are numbered: 2, 3, 4, 5, 6. Red arrows indicate schematic position of codons (numbers) that are tested. Historically uncoding exon 1 was referred to as 0; hence, rarely in literature data the naming of the exons is shifted. Due to alternative splicing, two isoforms of KRAS are produced. The isoforms differ at their 3’ ends: one contain exon 5 and the other exon 6, historically named 4A and 4B, respectively. Hence, protein isoforms are named: KRAS 4A (189 amino acids) or KRAS 4B (188 amino acids). The KRAS 4B, which accounts for 90–95% of all KRAS produced in cells, has anti-apoptotic functions and stimulates cell migration. On the contrary, KRAS 4A has pro-apoptotic properties [25–27]](image-url)
Although literature data indicate that mutations in \(BRAF\) and \(KRAS\) are mutually exclusive, there are sporadic reports of CRC tumours with coexisting \(KRAS\) and \(BRAF\) mutations [28–30]. Moreover, according to the recent American Society for Clinical Pathology, the College of American Pathologists, the Association for Molecular Pathology, and the American Society of Clinical Oncology guidelines, routine assessment of the \(BRAF\) status is only recommended for prognostic stratifications and in patients with deficient MMR tumours [31].

Nevertheless, there is a great deal of evidence that \(BRAF\) mutations may negatively affect anti-EGFR therapy. Therefore, local guidelines regarding the spectrum of molecular testing prior to anti-EGFR therapy may differ. For example, in Poland exclusion of \(KRAS\), \(NRAS\), and \(BRAF\) mutations is mandatory for anti-EGFR therapy consideration [32, 33].

Molecular diagnostic of these genes is often performed in separated tests, one after another: \(KRAS\), \(NRAS\), and \(BRAF\), according to mutation detection rate. If mutation is found in \(KRAS\), analysis of (an)other gene(s) is not performed (as patient is already disqualified from anti-EGFR therapy).

Novel findings indicate, however, that in some CRC tumours other \(BRAF\) mutations are detected, which lead to silencing of \(BRAF\) kinase activity. Such patients may respond to EGFR therapy. Importantly, if they also have an activating mutation in \(KRAS/NRAS\) genes, they may benefit from another type of targeted therapy – using MEK inhibitors. Thus, one can expect that within the next years \(BRAF\) testing may become compulsory regardless of \(KRAS/NRAS\) status [34].

Modern methods of mutation detection

Rapid development of molecular biology has enabled elaboration of many methods for DNA mutation detection (Table 3). The majority of them are based on PCR amplification of selected DNA fragments. The available methods, although widely used, differ both in the technical approach and sensitivity (defined as the minimal percentage of mutant cells that can be detected).

The simplest methods, usually qualitative ones, allow us only to determine if mutation(s) is/are present or not. More advanced techniques are based on modification of PCR reaction, require specialist equipment, and enable accurate determination of the mutation type. One of those techniques is direct sequencing, using Sanger method, which currently is called “a gold standard for mutation detection”. Furthermore, the other technique – pyrosequencing – enables determination of the amount of mutant allele in relation to the normal tissue. Modern, expensive techniques based on real-time PCR technology allow analysis of a large number of samples in a short time, while very sensitive and precise tests based on next-generation sequencing technology (NGS) additionally allow the simultaneous detection of many other molecular markers.

There are also CE-IVD certified kits available, which can be used to comply with the European Union directives. It should be remembered, however, that diagnostic CE-IVD tests are valid only when performed on CE-IVD equipment.

### Table 2. Mutations in the \(KRAS/NRAS/BRAF\) genes based on the COSMIC database

<table>
<thead>
<tr>
<th>Gene</th>
<th>Exon</th>
<th>Codon</th>
<th>Mutation name (protein level)</th>
<th>Mutation name (nucleotide level)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>KRAS</strong></td>
<td>2</td>
<td>12</td>
<td>p.Gly12Asp</td>
<td>c.35G &gt; A</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>p.Gly12Val</td>
<td>c.35G &gt; T</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>p.Gly12Cys</td>
<td>c.34G &gt; T</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>p.Gly12Arg</td>
<td>c.34G &gt; C</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>p.Gly12Ser</td>
<td>c.34G &gt; A</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>p.Gly12Ala</td>
<td>c.35G &gt; C</td>
</tr>
<tr>
<td></td>
<td>13</td>
<td>p.Gly13Asp</td>
<td>c.38G &gt; A</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>p.Gly13Cys</td>
<td>c.37G &gt; T</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>p.Gly13Arg</td>
<td>c.37G &gt; C</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>p.Gly13Ser</td>
<td>c.37G &gt; A</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>p.Gly13Ala</td>
<td>c.38G &gt; C</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>p.Gly13Val</td>
<td>c.38G &gt; T</td>
</tr>
<tr>
<td></td>
<td></td>
<td>59</td>
<td>p.Ala59Thr</td>
<td>c.175G &gt; A</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>p.Ala59Glu</td>
<td>c.176C &gt; A</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>p.Ala59Ala</td>
<td>c.176C &gt; G</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>59</td>
<td>p.Gln61Lys</td>
<td>c.181C &gt; A</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>p.Gln61Lys</td>
<td>c.180_181TC &gt; AA</td>
</tr>
<tr>
<td></td>
<td></td>
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<td>p.Gln61Arg</td>
<td>c.182A &gt; G</td>
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<td>c.183A &gt; C</td>
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<td>p.Lys117Asn</td>
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<td>c.437C &gt; T</td>
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<td><strong>BRAF</strong></td>
<td>15</td>
<td>600</td>
<td>p.Val600Glu</td>
<td>c.1799T &gt; A</td>
</tr>
</tbody>
</table>
Selected methods used in molecular diagnostics

The direct Sanger sequencing method is widely recognised as the “golden standard” in DNA mutation detection. The method can be used to determine all mutations in the tested genes, i.e. KRAS, NRAS, and BRAF, allowing for accurate mutation naming both at nucleotide and amino acid levels.

The advantage of this method is the detection of all possible changes in the analysed region of DNA, with the exception of large rearrangements of whole exon(s).

Specific mutations in KRAS/NRAS/BRAF genes can also be identified using commercial qPCR sets, which contain ready-to-use reagents, positive controls, and endogenous controls enabling PCR inhibitors detection. These kits identify the most frequent mutations in KRAS codons 12, 13, 59, 61, 117, and 146, and NRAS codons: 12, 13, 61, 117, and 146, and BRAF codon 600.

Importantly, the very high sensitivity of this type of method makes them especially prone to cross contamination of specimens. For example, if a microtome is not thoroughly cleaned after cutting a sample with a high mutation load, the next sample to be cut is at risk of being contaminated by remnants of the previous one, which can lead to false-positive results. To avoid this, the other, less sensitive method, i.e. Sanger sequencing, can be used. Nevertheless, when macrodissection is not possible, high-sensitivity tests may be highly beneficial.

The technological solutions for rapid, sensitive, and less-laborious detection of somatic mutations develop constantly. There is already a fully automated, CE IVD certified platform (Idylla, Biocartis) available. This device automatically isolates DNA form paraffin scraps and, subsequently, performs qPCR reactions, all within 4-5 hours. Nonetheless, despite automation, pathological examination prior to molecular testing is required, and only samples containing more than 10% of neoplastic fixation can be proceeded. According to its producer, the limit of detection (LOD) reaches 5% [40].

At the moment, the choice of diagnostic method depends on individual laboratory preferences. Nevertheless, regardless of the technology, thorough validation of the method it is crucial, which includes:
- definition of minimum amount of material and minimal thickness of paraffin scraps for DNA extraction,
- determination of the type of fixatives that can be used for biological material without the risk of negative impact on molecular procedures,
- definition of the quantity, quality, and concentration of the DNA sample that can be processed,
- determination of threshold value to distinguish the mutant from the normal variant,
- verification of test sensitivity (e.g. using dilution series),
- application of the reference method to verify the obtained results,
- ascertainment of method repeatability.

Furthermore, it is crucial to strictly adhere to the procedures, using positive and negative controls in each run and undergoing internal quality control. In addition, regular participation in external quality controls is equally important. Such international programs are annually carried out by EMQN (European Molecular Genetics Quality Network), UKNEQAS (United Kingdom National External Quality Assessment for Molecular Genetics), Colon EQA (European Society of Pathology in collaboration with the University of Leuven), and others.

According to recommendations, the sensitivity of the method should oscillate around 1-5% of tumour cells for allele-specific PCR methods and 10-20% for direct sequencing (Fig. 4). Regarding the specificity, at least seven typical changes in codons 12 and 13, variants in codons 59, 61, 117, and 146 of KRAS and NRAS, and variants occurring in codon 600 of the BRAF gene should be detected.

For the purpose of accreditation, the laboratory should obtain DNA in a minimum of 95% of cases and demonstrate correctness of analyses at the level of 97% [31, 36–38]. The acceptable turnaround time of the entire procedure should not exceed a maximum of 10 working days [36].

Although mutations detected using qPCR are automatically named, in the case of Sanger sequencing mutation, the name is usually assigned and verified by a diagnostician. Therefore, to avoid misinterpretation, it is extremely important to refer to a proper reference sequence (e.g.

### Table 3. Selected techniques used for KRAS, NRAS, and BRAF genotyping [13, 34–39]

<table>
<thead>
<tr>
<th>Technique</th>
<th>Sensitivity of the method*</th>
<th>Time of analysis</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sanger sequencing and pyrosequencing</td>
<td>10–20%</td>
<td>5–10 working days</td>
<td>Quite low sensitivity, the result requires interpretation by an experienced user, relatively low cost, the ability to detect new mutations, usually do not have CE IVD. This method is selected by approx. 20% of laboratories performing KRAS/NRAS/BRAF determinations</td>
</tr>
<tr>
<td>Methods based on allele-specific PCR – commercial kits</td>
<td>1–5%</td>
<td>Less than 5 working days</td>
<td>Reliable and repeatable techniques based on PCR, have high sensitivity and high specificity, the ability to detect specific mutations, require many molecular probes to be used, have CE IVD. This method is selected by approx. 50% of laboratories performing KRAS/NRAS/BRAF determinations</td>
</tr>
<tr>
<td>Next-generation sequencing (NGS) – commercial kits</td>
<td>0.1%-10%</td>
<td>Around 10 working days or more</td>
<td>Lack of full validation, expensive, TAT is usually longer due to the need of samples pooling, require interpretation by an experienced user, usually do not have CE IVD. This method is selected by approx. 30% of laboratories performing KRAS/NRAS/BRAF determinations</td>
</tr>
</tbody>
</table>

*The minimal content of cells with a mutation in relation to normal cells
Quality and practical aspects of pathological and molecular diagnostics in metastatic colorectal cancer (mCRC) according to LRG: https://www.lrg-sequence.org/) and to use standard mutation nomenclature as defined by HGVS (https://varnomen.hgvs.org/).

Proper and clear reporting of genetic test results is an extremely important issue, which in the case of somatic and tumour mutation diagnostics should be treated as a supplement to histopathological examination. Hence, information from pathological verification, concerning diagnosis and the percentage of neoplastic fixation in a specimen for molecular examination, is obligatory [37]. As stated above, pathological evaluation is crucial with respect to selection of the best methodology, and together with the reason for testing is indispensable for correct interpretation of the results. Consequently, according to European recommendations on mutation identification in KRAS, NRAS, and BRAF genes, the molecular test result should be submitted to the pathologist, who incorporates it into the full characteristics of the paraffin-embedded material or biopsy [31].

Alternative methods of KRAS/NRAS/BRAF testing

Analysis of circulating tumour DNA (ctDNA) is a relatively novel and valuable diagnostic method. In this strategy, mutation analysis is performed on DNA secreted by tumour cells into the bloodstream. Whenever a cancer sample cannot be obtained, the ctDNA analysis remains the only option. Mutation detection can be performed using ctDNA-dedicated high-sensitivity qPCR tests. Similarly, only dedicated tubes can be used for blood collection. Such tubes contain stabilisers preventing lysis of nucleated blood cells and the release of their genomic DNA, which, if present in the sample, would quantitatively dominate over ctDNA, and finally cause false results. Although ctDNA fraction is generally regarded as a valuable diagnostic material because it represents the entire tumour heterogeneity (not only its fragment, as in the case of standard FFPE tumour specimens testing), it is not recommended that the ctDNA be analysed when a tissue material is available. This is due to the fact that in some studies mutations detected in the tumour samples were not found at the ctDNA level [41].

Large-scale techniques such as next-generation sequencing (NGS) can also be used for genetic diagnostics in targeted therapy qualification analysis. The technology makes it possible to simultaneously analyse coding sequences of many genes, the entire exome, or even the whole-genome. Currently, in CRC only three molecular markers, KRAS, NRAS, and BRAF need to be evaluated by molecular testing; therefore, this laborious and still expensive approach is rarely used. Nevertheless, its sensitivity is very high – even less than 0.1% DNA with mutation in relation to the wild type DNA can be detected. This makes NGS a very good tool for tissue and ctDNA analysis. In the near future, NGS will probably become more popular because it will be routinely used to assess emerging genetic prognostic biomarker – tumour mutation burden (TMB), which plays a very important role in the immunological therapy of CRC [34].
Conclusions

Routine molecular diagnostics for targeted therapy is a relatively new discipline. Therefore, it is particularly important to develop standards and guidelines for material selection for molecular studies, and for mutation detection itself. In this initial phase, one should focus on a reliable examination of the basic prognostic factors for targeted therapy. In this initial phase, one should focus on a reliable selection for molecular studies, and for mutation detection a relatively new discipline. Therefore, it is particularly important to constantly search for new therapeutic goals, and thus to implement the widest possible panel of modern methods of molecular biology.

The authors declare no conflict of interest.


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