Clinical practice guidelines for the management of metastatic colorectal cancer: a consensus statement of the Hellenic Society of Medical Oncologists (HeSMO)

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Abstract

There is discrepancy and failure to adhere to current international guidelines for the management of metastatic colorectal cancer (CRC) in hospitals in Greece and Cyprus. The aim of the present document is to provide a consensus on the multidisciplinary management of metastatic CRC, considering both special characteristics of our Healthcare System and international guidelines. Following discussion and online communication among the members of an executive team chosen by the Hellenic Society of Medical Oncology (HeSMO), a consensus for metastatic CRC disease was developed. Statements were subjected to the Delphi methodology on two voting rounds by invited multidisciplinary international experts on CRC. Statements reaching level of agreement by ≥80% were considered as having achieved large consensus, whereas statements reaching 60-80% moderate consensus. One hundred and nine statements were developed. Ninety experts voted for those statements. The median rate of abstain per statement was 18.5% (range: 0-54%). In the end of the process, all statements achieved a large consensus. The importance of centralization, care by a multidisciplinary team, adherence to guidelines, and personalization is emphasized. R0 resection is the only intervention that may offer substantial improvement in the oncological outcomes.

Keywords Colorectal cancer, metastatic disease, surgery, guidelines

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Introduction

Factors predisposing to metastases

An increased risk of metastases is anticipated in case of: 1) advanced primary disease (advanced T stage, positive lymph nodes, presence of extramural venous invasion, poor differentiation) [1-3]; 2) female gender [3]; 3) location at the rectum [3]; or at left than the right colon [4]; 4) increased CD-10 expression [5]; 5) KRAS and PIK3CA
bi-mutations; 6) presence of metabolic syndrome [6]; and 7) increased serum levels of carcinoembryonic antigen (CEA) (>5 ng/mL) alanine aminotransferase, aspartate aminotransferase, glutamyltransferase, Lactate Dehydrogenase (LDH), Vascular Endothelial Growth Factor Receptor (VEGFR) and Tissue Inhibitor of Metalloproteinases 1 (TIMP-1) [1,7]. In contrast, patients with hepatitis B virus infection present a reduced risk of liver metastasis and increased rate of R0 resections, but also increased risk of extra-hepatic metastasis [8]. There is some evidence that portal vein chemotherapy with 5-fluorouracil (5-FU) could be an effective preventive measure for liver metastases. According to a recent study [9], 5-year liver metastasis rate is 12.5% after curative surgery for the primary tumor combined with portal vein chemotherapy, opposed to a 25% relative rate after curative surgery alone.

**Aim**

The Hellenic Society of Medical Oncology (HeSMO) selected an executive team on the grounds of their experience in colorectal cancer (CRC) and hepato-biliary and pancreatic malignancies which was assigned to develop a consensus statement and form guidelines on the main aspects of image staging, surgical treatment and follow up of metastatic CRC, based on the review of literature and the principles of the evidence-based medicine.

The present draft is part of a large consensus on the guidelines for the management of colorectal cancer. Guidelines on: 1) epidemiology, molecular biology, genetics, prognostic and predictive markers, hereditary forms, surveillance; 2) colon cancer care; 3) rectal cancer care; and 4) adjuvant treatment of CRC are presented elsewhere.

**Methodology**

The methodology in setting our guidelines for the surgical management of rectal cancer has already been reported elsewhere [10]. The first round of the online Delphi voting process opened on September 29th 2013 and closed on December 6th 2013. The second round opened on January 6th 2014 and closed on January 24th 2014. In the final document, all statements are presented as recommendations of care. Even statements achieving a consensus of <80% were included. At the end of each recommendation the level of evidence (LOE) and the strength of recommendation (SOR) are mentioned, followed by the rate of voting consensus (ROVC) (Table 1).

**Legal disclaimer**

Details on the legal aspects of these guidelines have already been reported [10].

**Discussion**

Ninety experts participated in the Delphi methodology and voted for 101 statements. Median abstain rate was 18.5% (0-54%). After the first voting process, 24 statements achieved a 100% voting consensus by all participants. There were 61 statements achieving a 90-99% consensus, and 13 achieving a consensus rate of 80-89%. Three statements achieved a rate of consensus of less than 80%. These three statements were circulated among the members of the executive team and, after being amended, they re-entered a second round of voting. Also, seven new statements entered directly the second voting round making the total number of statements 108. At the end of the process, there were no statements with a ROVC of less than 80% (Table 2).

**Table 1 Evidence level and recommendation grade**

| Level of evidence | Description |
|-------------------|-------------|
| I                 | Evidence from at least one large RCT of good methodological quality (low potential for bias) or meta-analyses of well-conducted RCTs without heterogeneity |
| II                | Small RCTs or large RCTs with a suspicion of bias (lower methodological quality) or meta-analyses of such trials or of trials with demonstrated heterogeneity |
| III               | Prospective cohort studies |
| IV                | Retrospective cohort studies or case-control studies |
| V                 | Studies without control group, case reports, experts opinions |

**Strength of recommendation**

| Grade | Description |
|-------|-------------|
| A     | Strong evidence for efficacy with a substantial clinical benefit, strongly recommended |
| B     | Strong or moderate evidence for efficacy, but with a limited clinical benefit, generally recommended |
| C     | Insufficient evidence for efficacy or benefit does not outweigh the risk or the disadvantages (adverse events, costs) optional |
| D     | Moderate evidence against efficacy or for adverse outcome, generally not recommended |
| E     | Strong evidence against efficacy or for adverse outcome, never recommended |

**Table 2 Rate of voting consensus of statements after the two voting processes**

| Rates of voting consensus | Statement numbers after 1st voting process | Resubmitted statement numbers | Statement numbers at the end of process |
|--------------------------|------------------------------------------|-------------------------------|----------------------------------------|
| 100%                     | 24                                       | 24                            |                                        |
| 90-99%                   | 61                                       | 69                            |                                        |
| 80-89%                   | 13                                       | 15                            |                                        |
| 70-79%                   | 1                                        | 1                             |                                        |
| 60-69%                   | 2                                        | 2                             |                                        |
| New statements           | 7                                        |                               |                                        |
| Total: 101               | Total: 10                                | Total: 108                    |                                        |

**RCT, randomized control trial**
General considerations

Background

All patients with metastatic CRC, and in particular candidates for resection, should be discussed and managed in the context of a multidisciplinary team (MDT) [11]. Such a team should comprise the necessary medical and non-medical team members (Diagnostic: radiologists, pathologists; Therapeutic: colorectal, hepatobiliary and thoracic surgeons, medical oncologists, radiotherapist, palliative care medical personnel; and Others: nursing). Treatment of metastatic CRC should be personalized and based on multimodality measures. Also, it has been documented that liver surgery in high-volume centers, practiced by highly specialized and experienced personnel, is associated with better outcomes including morbidity, perioperative mortality and survival [12-14].

RECOMMENDATIONS

1. Treatment should be personalized, based on multimodality measures and after MDT meeting discussions (SOR A) (ROVC: 100%)  
2. Centralization of management offers the best short-term outcomes, namely mortality of 5% and 90-day morbidity of 30-40% after hepatic resection (SOR A) (ROVC: 97%)

Imaging of metastatic disease – other staging modalities

Background

Chest x-ray may be the initial approach to detect pulmonary metastasis, but in general thoracic and abdominal computed tomography (CT) is recommended to detect or rule out distant metastases. The real value of CT is its accuracy in detecting distant metastases. Magnetic resonance imaging (MRI) is helpful in further characterization of liver lesions suspected for metastases diagnosed by CT scan or ultrasound (US) [15]. MRI is the preferred first-line modality for evaluating colorectal liver metastases in patients who have previously undergone treatment, and for resolving diagnostic problems with equivocal lesions. In a recent meta-analysis [16], MRI sensitivity in detection of colorectal metastases was higher than of CT, particularly for small lesions (<10 mm). In an even more recent report enhanced MRI was proven more accurate than multi-detector CT (MDCT) or US for the evaluation of liver metastases [15]. Enhanced MRI with diffusion-weighted series is more sensitive and accurate in detecting CRC liver metastases than the non-enhanced MRI [17].

Fluorodeoxyglucose-positron emission tomography (FDG-PET) could be considered for detection of liver metastases and peritoneal disease when there is clinical, biochemical or radiological suspicion of systemic disease [18]. FDG-PET is mainly useful in the assessment of local recurrence and metastatic disease, when conventional imaging is not helpful [19]. Furthermore, PET/CT is more accurate than CT alone in detection of metastatic disease, in patients with high suspicion for recurrence [20]. Also, PET/CT is more reliable than MDCT for the detection of small subcapsular liver metastatic deposits [21]. Currently, FDG-PET is not used as a primary staging modality in CRC. Bone scan and brain imaging are only helpful in case of relevant clinical symptoms.

There is a limited benefit from laparoscopic staging of CRC metastatic disease with the use of US. According to a recent meta-analysis [22], laparoscopic US staging has a low sensitivity of 59%, and is only recommended as an additional diagnostic modality to suspected peritoneal carcinomatosis. None of the available imaging or intraoperative diagnostic modalities is sensitive or accurate in the assessment of the lymph nodes at the hepatic hilum [23].

RECOMMENDATIONS

3. Abdominal CT or MRI is required to detect liver metastases (LOE V; SOR B) (ROVC: 98%)  
4. US is less accurate than MRI in the detection of liver metastases (LOE I, SOR A) (ROVC: 95%)  
5. Chest X-ray and, preferably chest CT is recommended for the detection of pulmonary metastases (LOE V; SOR B) (ROVC: 95%)  
6. MRI is the preferred imaging modality for the evaluation of previously treated colorectal cancer on equivocal liver metastases (LOE IV; SOR B) (ROVC: 89%)  
7. FDG-PET should not be routinely used for the detection of metastatic disease, unless there is biochemical evidence of recurrent disease in the absence of apparent lesion (LOE V; SOR D) (ROVC: 100%)  
8. Bone scan and brain imaging should only be performed for patients presenting relevant symptoms (LOE V; SOR D) (ROVC: 97%)  
9. Laparoscopic staging of liver metastasis with the use of US is of limited accuracy (LOE I, SOR A) (ROVC: 84%)

Management of resectable liver metastases (Fig. 1)

Indications for surgical resection

Background

R0 resection is the treatment of choice for the CRC metastatic liver disease. For patients with resectable liver metastases surgical resection offers the best chance for long-term survival with actuarial 5-year survival rates ranging from 30% to 58% in some selected series, and a cure rate of 20% [24-26]. However, disease relapses, mostly in the liver, in 60-75% of these patients. In general, criteria for resectability of liver or lung metastases are not standardized and have evolved over the last years. They are clearly related to the experience of the surgeon and the
Figure 1

Staging of Metastatic Disease
MDT Discussion

- R0 Resection Feasible
  - Single Liver Metastasis <2cm
  - Single Liver Metastasis >2cm or Multiple Liver Metastases

- R0 Resection Questionable
  - Up-Front CTx
    - FOLOX 3m
  - Intensive CTx
    - FOLOX 3-6m +/-
    - Other Measures
      - PV Ligation
      - PV Embolisation
      - HAI

- R0 Resection Not Feasible
  - Intensive CTx
    - Escalation
    - Change

- Resection
  - Adjuvant CTx
    - up front no: FOLOX 6m
    - up front yes: FOLOX 3m

MDT assessment. Based on data from old series, unresectable disease is considered in the presence of: 1) more than four metastatic lesions; 2) bilobar disease; 3) extrahepatic disease; and 4) inability to achieve resection margins of at least 1 cm [27,28]. None of these factors is considered now as absolute contraindication for surgery; however, patients with these characteristics are those who have a worse prognosis. Current indications for resection of CRC liver metastases, are removal of all deposits with adequate clear margin, and preservation of sufficiently function of hepatic parenchyma. A prerequisite for resection of liver metastases is a functioning remnant of at least 30% of the total volume of the liver. Resectional surgery should...
not endanger patient’s life because of unsuitable location of metastases. The ability to achieve R0 resection and assessment of the volume of healthy liver remnant should be determined by the radiologist and surgeon preoperatively. MDCT-volumetry is considered as the method of choice to estimate remaining liver volume [29-31]. Furthermore, MR spectroscopy can be applied for the assessment of liver steatosis in patients who have received neo-adjuvant chemotherapy [32]. Additional considerations, prior to deciding resection of liver metastases, are any possible extrahepatic recurrence and poor tumor biology, as these factors adversely affect outcomes.

RECOMMENDATIONS

10. Hepatic R0 resection is the treatment of choice for liver metastases, associated with a 5-year survival rate of 58% and a cure rate of 20% (LOE II; SOR A) (ROVC: 99%)

11. Survival after systemic chemotherapy followed by surgery is superior to chemotherapy alone (LOE III; SOR B) (ROVC: 93%)

12. Age, number and location of lesions and extrahepatic disease (if resectable) are not contraindications for surgery (LOE III; SOR C) (ROVC: 87%)

13. The aim of resection of liver metastases is to remove all macroscopic disease with clear (negative) margins (LOE II; SOR B) (ROVC: 100%)

14. The ability to achieve clear margins (R0 resection) and assessment of the volume of healthy liver remnant should be determined by the radiologist and surgeon preoperatively (LOE III; SOR C) (ROVC: 100%)

15. MDCT volumetry is considered as the method of choice to estimate remaining liver volume (LOE III; SOR C) (ROVC: 99%)

16. Intraoperative US is the only method capable to detect the smallest liver metastases that may be missed by other imaging modalities (LOE III; SOR A) (ROVC: 93%)

17. At one stage surgery, multiple resections can be performed, provided there is sufficient remnant liver (>30%) and surgery is not too risky due to the location of the lesions. Other considerations must include the presence of questionably resectable extrahepatic disease and poor tumor biology (LOE IV; SOR A) (ROVC: 97%)

18. There is a tendency towards segmental or wedge resections, instead of major hepatectomy (LOE III, SOR B) (ROVC: 96%)

Perioperative chemotherapy

Background

The role of perioperative chemotherapy in resectable CRC metastatic liver disease is unclear, particularly in case of a solitary, <5 cm metastatic lesion. For resectable multiple metastatic disease of the liver, an EORTC trial using perioperative chemotherapy with FOLFOX (3 months before and 3 months after metastasectomy) showed superior progression-free survival (PFS) for the perioperative treatment group [33]. In addition, some more recent studies [34,35] have shown a survival benefit with perioperative chemotherapy. Although final survival data are still awaited, this approach represents a current standard with level 2 evidence. There is no evidence yet that adding a biologic to a cytotoxic doublet may improve outcome in resectable metastases, compared with a cytotoxic doublet alone. Intensification of induction chemotherapy is currently under investigation (FOLFOX beva/anti-endothelial growth factor receptor (anti-EGFR), FOLFIRI+/-beva/anti-EGFR). The optimal sequencing of chemotherapy in patients with resectable liver metastases (perioperative versus postoperative) was the subject of an ongoing NCI-sponsored trial (NSABP-C-11) but the study has been closed prematurely due to low accrual.

For patients with resectable disease and insufficient response on preoperative chemotherapy immediate metastasectomy may be considered. In case of progressive disease during preoperative chemotherapy, second line systemic chemotherapy should be administered. Alternatively, hepatic intra-arterial (HIA) chemotherapy in combination with systemic chemotherapy can be considered in very experienced centers.

RECOMMENDATIONS

19. For patients with primarily resectable lesions, treatment with FOLFOX for 3 months preoperatively, followed by resection, followed by 3-month FOLFOX treatment could be considered (LOE II; SOR A) (ROVC: 82%)

20. In case of solitary metastatic tumor <5 cm, use of pre-operative chemotherapy is unclear (LOE IV; SOR B) (ROVC: 97%)

21. During preoperative treatment, a thorough evaluation for complete remission has to be performed, after 6 weeks of treatment (LOE III; SOR B) (ROVC: 93%)

22. If preoperative chemotherapy was not applied, in case of R0 resection, adjuvant chemotherapy, as in stage III colon cancer, for 6 months can be considered (LOE III; SOR A) (ROVC: 99%)

23. Timing interval from chemotherapy to surgery should be 4-6 weeks, as also from surgery to adjuvant chemotherapy (LOE I; SOR A) (ROVC: 96%)

24. For patients with resectable disease and insufficient response on preoperative chemotherapy immediate metastasectomy may be considered (LOE III; SOR B) (ROVC: 97%)

25. In case of disease progression during preoperative chemotherapy second-line systemic chemotherapy should be administered (LOE II; SOR A) (ROVC: 94%)

26. Alternative option for progressive disease is the use of HIA plus systemic chemotherapy, but only at very experienced institutions (LOE II; SOR B) (ROVC: 96%)
Surgery

Background

An intraoperative staging is always recommended with the use of standard and contrast enhanced US. Intraoperative US may identify small solitary or multiple liver metastatic lesions, not preoperatively detected by other imaging modalities. In a recent study [36], 18% additional lesions were identified in 44% of the patients. At surgery for the resection of CRC liver metastases, concomitant resection of the hepatic hilar lymph nodes is recommended for staging reasons [37]. Positive hilar lymph nodes can be found in up to 11.5% of patients, and may be associated with unresectability of the liver deposits in 28.6% of the cases [38]. Concomitant resection of involved lymph nodes may improve survival [37].

The standard procedure for liver metastases is a R0 resection of all lesions, sparing at the same time as much hepatic parenchyma as possible. For this reason, there is a tendency to shift from a typical liver lobectomy to multiple segmental or wedge excisions of metastatic lesions with adequate clear margin [39,40].

In case of multiple resectable metastatic deposits the resection of which may not leave sufficiently functioning liver parenchyma, the two-stage surgery is recommended according to the following sequence: 1) 4-6 cycles of chemotherapy followed by imaging assessment; 2) surgery with wedge limited resections at the lobe which is to remain in situ complemented with ligation (PVL) or embolization of the portal vein (PVE) branch of the lobe to be removed; 3) optional chemotherapy and imaging reassessment after 4-6 weeks; 4) hepatectomy, provided that the remaining lobe is assessed as functionally sufficient; and 5) chemotherapy [41-44]. It has been suggested that chemotherapy, including targeted agents, at the interval between the two stages, does not impair liver degeneration after PVL [45,46] and that omission of interval chemotherapy may result in progression of liver disease [47]. The choice of one- over two- stage surgery for resectable multiple metastatic lesions should be personalized according to tumor burden, anatomical location of deposits, anticipated functional sufficiency of hepatic remnant and presence of extrahepatic disease. It should be considered that resection of >4 lesions at one stage, is associated with less R0 resections and recurrence rates that exceed 58% [48]. Presence of primary colorectal lesion and progression of the disease after the first stage preclude two-stage liver surgery [44].

The in situ split procedure (PVL of the intended hepatic lobe to be resected and split of the hepatic parenchyma at the intended transection line) aims to induce rapid remnant liver parenchyma hypertrophy, allowing hepatic lobectomy at a second stage during the same admission. However, because the reported mortality is high and available evidence very limited the procedure cannot be recommended unless preformed in highly specialized centers and within the frame of research trials.

Specific technical considerations

According to a meta-analysis [49] and a more recent study [50], application of the Pringle maneuver (temporary clamping of the porta hepatitis during hepatic resection) does not seem to have any effect on recurrence rate, the disease-free survival (DFS) or the overall survival (OS).

The laparoscopic approach for the resection of CRC liver metastases, has been applied in small series, by highly specialized surgical personnel. Immediate postoperative morbidity and mid-term survival are similar, while blood loss is less and length of hospital stay is shorter compared to the open approach [51]. There are also limited data on the robotic approach for the resection of CRC liver metastases from specialized centers [52].

RECOMMENDATIONS

27. After down-sizing or after complete response to preoperative chemotherapy, resection should include all initial sites of disease, if possible (LOE III; SOR A) (ROVC: 99%)

28. Removal of some and not all lesions after preoperative chemotherapy is not recommended (LOE II; SOR B) (ROVC: 99%)

29. In case of multiple bilateral lesions not amenable to one-stage surgery, two-stage surgery should be offered as follows: 1) 4-6 cycles of systemic chemotherapy and then imaging reassessment, followed by 2) surgery with small resections at the lobe to remain (if remnant is insufficient: PVL or PVE of the branch of the lobe to be resected), followed by 3) optionally systemic chemotherapy and imaging reassessment after 4-6 weeks, followed by 4) hepatectomy, followed by 5) systemic chemotherapy (LOE III, SOR B) (ROVC: 97%)

30. Neither chemotherapy nor bevacizumab interfere with liver regeneration, when given at the interval between the two stages (LOE II, SOR B) (ROVC: 89%)

31. Despite the potential liver damage, omission of interval chemotherapy between the two stages results in tumor cell proliferation (LOE III, SOR B) (ROVC: 92%)

32. Choice of one- over two-stage surgery is personalized according to tumor burden, anatomical tumor location, potential R0 resection, sufficient liver remnant and extrahepatic disease (LOE III, SOR B) (ROVC: 100%)

33. In case of more than 4 metastatic lesions, one-stage surgery is associated with less R0 resections and more recurrences (LOE III, SOR B) (ROVC: 85%)

34. Because mortality is high and available evidence very limited, the in situ split staged procedure cannot be recommended unless preformed in highly specialized centers and within the frame of research trials (LOE IV; SOR D) (ROVC: 100%)

35. Due to lack of high level of evidence, the laparoscopic approach for the resection of hepatic metastases is not recommended, unless performed by a very experienced surgical team and within the context of clinical trials (LOE IV; SOR D) (ROVC: 97%)
Ablation

Background

In recent years, there is a tendency towards limited resections in combination with ablation of small lesions, instead of surgery alone for the treatment of multiple resectable metastatic lesions, aiming at preserving hepatic parenchyma [53]. Combination of surgery with ablation is indicated in the presence of increased metastatic burden, when complete surgical resection is not feasible and when ablation is applicable to small <3 cm lesions not in the vicinity of major vessels. Oncological outcomes after surgery alone is superior to the combination of surgery with ablation, but the latter approach is applied to patients with increased metastatic burden [54-57]. In general, ablation either by radiofrequency (RFA) or microwaves (MWA) as a sole treatment is inferior to surgery, and apart as an adjunct to surgery, is reserved for patients unfit to undergo surgery or as a bridge to surgery [29].

There has been a debate whether a solitary liver metastasis could be treated by resection or ablation with the use of radiofrequency. According to a meta-analysis [58] and some additional comparative studies [59-61] resection of the solitary metastatic lesion, particularly when >3 cm, is associated with better oncological outcomes (OS and DFS) compared to RFA. For hepatic metastatic lesions <3cm both modalities offer similar OS and DFS [59].

RECOMMENDATIONS
36. Ablation is generally inferior to resection. However, it can be used, preferably with the use of RFA or MWA: 1) in patients unfit for surgery; or 2) as bridge to resection; or 3) in combination with resection (if liver remnant is not considered functionally sufficient) (LOE IV; SOR D) (ROVC: 100%)
37. Surgery combined with ablation is indicated: 1) in case of increased metastatic burden; 2) when complete resection is not feasible; and 3) when small lesions (<3 cm) to be ablated are not in the vicinity of major vessels (LOE II, SOR B) (ROVC: 99%)
38. Surgery, compared to surgery combined with ablation, offers better oncological outcomes, although the latter approach is reserved for patients with greater tumor burden (LOE II, SOR B) (ROVC: 99%)

Adjuvant chemotherapy

Background

The role of adjuvant chemotherapy after R0 metastasectomy is still unclear. Also, unclear is the role and efficacy of chemotherapeutic regimes including targeting agents. Underpowered trials with 5-FU and levamizole (LV) or HIA with FUDR+ systemic chemotherapy with 5-FU+LV have shown some advantage, although no study has shown a clearly significant survival benefit [62,63]. Nevertheless, postoperative adjuvant systemic chemotherapy is recommended, as in the case of stage III colon cancer, in Europe and USA, despite lack of level 1 or 2 evidence. In an ongoing phase II/III trial in Japan, adjuvant FOLFOX6 was compared to surgery alone [64].

Currently the value of bevacizumab, in addition to adjuvant capecitabine (XEL) and oxaliplatin (XELOX), after liver metastasectomy, is being investigated in the Netherlands. Another trial (NSABP C-09), comparing adjuvant systemic oxaliplatin plus XEL alone or with HIA FUDR, has been closed prematurely due to the low accrual.

RECOMMENDATION
39. There is no substantial evidence that adjuvant chemotherapy after R0 hepatic metastasectomy offers to oncological outcomes (LOE IV, SOR B) (ROVC: 91%)

Other adjuvant therapeutic modalities

Background

Initially, there has been evidence that administration of HIA in addition to systemic chemotherapy after R0 resection of liver metastasis offers better 2-year survival and longer PFS than systemic chemotherapy alone [65,66]. However, a systematic review [67] showed that HIA has no additional benefit to adjuvant systemic chemotherapy on survival.

RECOMMENDATION
40. HIA as an adjunctive modality after R0 hepatic metastasectomy is of no benefit (LOE III, SOR B) (ROVC: 96%)

Immediate postoperative outcomes

Provided that patients are treated in high volume centers, with highly specialized and experienced personnel and have been subjected to thorough preoperative assessment, immediate postoperative morbidity and mortality are low. Morbidity and mortality depend on several parameters, which include extent of perioperative chemotherapy, amount of resected hepatic parenchyma, loss of blood and needs for blood transfusion [68-70].

Postoperative liver failure is seen in approximately 10% of cases and depends on the functional adequacy of the remaining hepatic parenchyma. Postoperative liver failure has a remarkable impact on postoperative mortality [69].
Management of resectable lung metastases

Background

Patients with limited lung metastases should be subjected to surgical R0 resection. Resection of lung metastases can be performed by thoracoscopy with equal or even better short-term results than after thoracotomy. The 5-year survival rate after R0 resection is 35-50%, and depends on multiplicity and size of metastatic lesions, status of hilar or mediastinal lymph nodes, and history of liver metastasis [71-74]. The 5-year recurrence rate after the first metastasectomy is 35% by thoracoscopy and 21% by thoracotomy [74].

Despite the lack of data regarding perioperative treatment, an approach similar to the management of resectable liver metastases should be considered. A FOLFOX regimen is recommended for 3 months prior and 3 months after resection mainly based on the data from stage III disease and those for the perioperative treatment of after liver resection [33]. Alternatively, an initial resection, without perioperative treatment, followed by postoperative chemotherapy can be applied.

RECOMMENDATIONS
41. Successful resection of lung metastases, offers a 5-year survival rate in almost one third of patients (LOE III; SOR B) (ROVC: 100%)
42. Peri- or post-operative chemotherapy is recommended (LOE II; SOR A) (ROVC: 95%)

Treatment of advanced metastatic disease (Fig. 2)

The aim of the first-line treatment depends on the clinical presentation and biology of the tumor (metastases limited to liver or lung or both, or peritoneum; dynamics of progression; symptoms) and patient factors (co-morbidities, age, potential to undergo secondary resection). In case of major response to induction chemotherapy of liver, lung (or even peritoneal) R0/R1 resection can result

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**Figure 2** Strategy and treatment algorithm of metachronous advanced metastases (liver +/- lungs)
MDT, multi-disciplinary team; CTx, chemotherapy; PV, portal vein; HAI, hepatic arterial infusion chemotherapy; FOLOX, 5-fluoro-uracil plus oxaliplatin
in long-term survival and potentially cure in a minority of patients. With this respect, patients can be individually categorized into the following three clinical groups, by parameters describing localization and extent of disease, tumor dynamics, co-morbidities, potential of the patient to tolerate chemotherapy and surgical treatment: 1) liver (± lung) metastases only, which may become resectable after induction chemotherapy ± limited/localized metastases to other sites, and subject physically able to undergo major surgery (biological age, heart/lung condition); 2) multiple metastases with rapid progression, tumor-related symptoms and risk of rapid deterioration, with co-morbidity allowing aggressive treatment; and 3) unresectable metastases – never suitable for resection, without major symptoms or risk of rapid deterioration, or with severe co-morbidity, precluding surgery and/or intensive systemic treatment.

Molecular factors

With the exception of RAS mutation, which precludes patients from treatment with anti-EGFR antibodies, there is no other prognostic/predictive molecular marker, relevant to routine first-line treatment, which affects decision outside clinical trials in advanced metastatic disease [75-78].

Clinical factors

Background

Although never prospectively proven, the achievement of a disease-free status after chemotherapy and surgery, is the only mean to give the potential for long-term survival or cure, in an otherwise incurable/palliative situation. For this aim, the most active induction chemotherapy which is able to induce downsizing as much as possible in as many patients as possible, should be selected for group I. Furthermore, downsizing after chemotherapy is a marker of favorable tumor behavior [42,79-82]. A chemotherapeutic scheme of 5-FU, oxaliplatin and irinotecan succeeded to convert initially unresectable to resectable liver metastases, in a limited cohort of patients [83]. For the intermediate group II, the aim is a rapid regression of metastases, particularly in cases of imminent tumor associated complications. An escalation strategy, namely single agent chemotherapy followed by combination, could be hazardous because, at the time of switch to a more effective second-line treatment, it might be “too little - too late” for a favorable outcome. Patients of group iii are treated mostly with a palliative intent. Therefore, an escalation strategy seems appropriate. However, a small percentage of these patients with excellent responses might also become candidates for secondary surgery with further escalation [84].

Choice of chemotherapeutic regimen

Background

Available chemotherapeutic agents are fluoropyrimidines (5-FU modulated by folinic acid [FA], or XEL), irinotecan and oxaliplatin. XEL can be safely substituted by 5-FU/FA in combination with oxaliplatin (FOLFOX) and in combination with irinotecan (FOLFIRI), after the necessary dose adjustment of both drugs. However, FOLFIRI should be the preferred option over XELIRI as it is associated with less toxicity from the gastrointestinal tract [85].

Monoclonal antibodies are the vascular endothelial growth factor (VEGF) targeting bevacizumab and the anti-EGFR antibodies cetuximab and panitumumab. Anti-EGFR antibodies have no activity in RAS mutant tumors. The combination of XEL, oxaliplatin with cetuximab, and also panitumumab, seem to have no additional benefit over XEL and oxaliplatin alone and should be avoided at present [86].

Standard combination chemotherapy regimens comprising 5-FU/LV in combination with either irinotecan, typically FOLFIRI, or oxaliplatin (FOLFOX) have been reported to facilitate the resection of 7-40% of patients with initially unresectable metastases depending upon the initial selection of patients [33]. However, 75-80% of these patients experience cancer relapse within 2 years of resection. Data emerging from randomized trials suggest that the addition of a targeted agent (bevacizumab or cetuximab) or even scarce data of phase II trials on the combination with a third cytotoxic plus or minus a targeted agent, might be even more effective, although concerns about toxicity limit the use of this triplet to highly selected cases. The combination of a chemo-doublet plus cetuximab has led to higher resection rates (although still low in absolute numbers) in patients with liver limited unresectable metastatic RAS wild-type (wt) CRC. The combination of FOLFOX/cetuximab and FOLFIRI/cetuximab has led to similar response rates and resection rates in RAS wt tumors [87,88].

The combination of a fluoropyrimidine/oxaliplatin/bevacizumab has led to a non-significant trend in an increased resection rate compared with the chemo-backbone alone, although no increase in response rate was shown [89]. In a more recent study [90], it was shown that backbone chemotherapy combined with bevacizumab is associated with increased
response and resectability rates and improved survival. There are no data available from randomized studies comparing a chemo-doublet plus bevacizumab with a chemo-doublet plus cetuximab.

Limited data exist concerning the three-drug combination FOLFOXIRI and the possibility to use it as an alternative option to FOLFIRI/FOLFOX + cetuximab with probably more toxicity, especially when targeting agents such as cetuximab are not available or are contraindicated [91,92]. Possible chemotherapeutic regimens, also including targeting agents are listed in Table 2.

First-line regimens depend on the chosen treatment strategy. For potentially resectable and/or symptomatic disease, first-line treatment should ideally be a chemotherapy doublet in combination with a monoclonal antibody or a triplet. If tumor shrinkage is desirable, treatment with FOLFIRI and cetuximab for RAS wt or FOLFOXIRI should be considered [93]. First-line treatment with monotherapy [94] or doublet (either chemotherapy or chemotherapy with monoclonal antibodies) could be a valid option for patients, for whom a second resection is not feasible, and who have no symptoms or risk for deterioration of the disease [95].

Selected induction chemotherapy should be given for at least 8 weeks, and first re-assessment is performed, in order to avoid unnecessary chemotherapy application in case of early progression [96,97]. If the treatment aim is pure palliation, the timing of first evaluation is of less importance, and an interval of 12 weeks might be more appropriate [98]. The overall induction treatment duration depends on the treatment aim. If secondary surgery is attempted, induction chemotherapy should be continued until R0 resectability can be achieved, with the first evaluation after 8 weeks, as mentioned above. If the chosen chemotherapeutic regimen continues to be active, a 4-month duration of treatment is recommended; if resectability is still not achievable, duration of treatment can be extended to 6 months. Further treatment with the same regimen is not recommended [98].

If the aim of secondary resection cannot be achieved, as well as in patients where resection is not the aim of treatment, this should be continued according to the individual's situation, patients' needs, cumulative toxicity, particularly with oxaliplatin and aggressiveness of the disease [99].

In most patients, survival is not significantly affected, if first-line treatment is not given continuously until progression, particularly in case of oxaliplatin discontinuation because of cumulative neurotoxicity [100]. Drug and treatment “holidays” are acceptable options in selected patients, after a substantial initial period of treatment [100,101]. Recently presented data support that de-escalation of chemotherapy combined with continuation of bevacizumab after induction treatment withXELOX + bevacizumab lead to significant benefit in PFS, but with limited impact in median OS [95].

The selection of the subsequent treatment lines should be based on the type of first-line treatment, response and/or PFS to first-line treatment, patient's performance status (PS) and preferences and taking into consideration the cost of treatment. New agents such as aflibercept (in second-line treatment) or regorafenib (≥ third-line treatment) could be considered in specific patient subpopulations.

**RECOMMENDATIONS**

46. Aggressive treatment regimens are FOLFOXIRI (LOE II; SOR C), FOLFIRI (LOE I; SOR B) and FOLFOX (LOE I; SOR B) and could be used alone or with the addition of anti-EGFR antibodies (RAS wt) or bevacizumab (LOE II, SOR B) (ROVC: 100%)

47. Duration of induction treatment may exceed 6 months, in case of favorable response (LOE II; SOR B) (ROVC: 84%)

48. RAS mutation precludes patients from treatment with anti-EGFR antibodies at the preoperative settings (LOE II, SOR B) (ROVC: 100%)

49. Alternative regimes, in patients with progressive metastatic disease are mXLIRI+/-bevacizumab or cetuximab (RAS wt) (LOE II; SOR C), or FOLFOX + panitumumab/ cetuximab (RAS wt) (LOE II; SOR B) (ROVC: 100%)

50. For RAS wt tumors, induction treatment with FOLFIRI + cetuximab or FOLFOX + anti-EGFR antibodies appears to be more effective in terms of major tumor shrinkage and secondary resectability, than bevacizumab based combinations, for which less data are available (LOE II; SOR B) (ROVC: 89%)

51. FOLFOXIRI should be considered as a treatment option especially for patients with RAS mutant tumors (LOE II; SOR C) (ROVC: 96%)

52. For palliation: i) fluoropyrimidine ±bevacizumab (sequential treatment) (LOE II; SOR B); or ii) doublets chemotherapy (LOE I; SOR A); or iii) doublets chemotherapy + anti-EGFR (RAS wt) (LOE II; SOR B) can be given (ROVC: 100%)

**Additional upfront measures**

**Background**

HIA can be used to downsize hepatic metastases and render them resectable, with improvement in survival rates [102,103]. This modality can be applied either as a first-line neo-adjuvant treatment in combination with systemic chemotherapy [104], or as a second-line approach in combination with chemotherapy when first-line systemic chemotherapy has failed to downsize metastatic lesions. Still, effectiveness of this modality remains questionable [105,106].

The initial chemotherapeutic regimen contained floxuridine and dexamethasone; the latter was given for the prevention of hepatic and biliary toxicity [107-110]. Other combined agents, such as 5-FU and leucovorin with or without oxaliplatin, have been given to reduce hepato-biliary toxicity and improve oncological outcomes, but data are limited and evidence is low at present [105,110-114]. HIA chemotherapy with novel agents
is tested, aiming at improvement of oncological outcomes. It is evident that more trials are needed to assess any additional survival benefit of HIA to systemic chemotherapy, including new chemotherapeutic regimens.

**RECOMMENDATIONS**

53. The rationale of HIA modalities is that: 1) hepatic metastases are primarily supplied by branches of the hepatic artery; and 2) larger quantities of the therapeutic agent in use are delivered to tumor site and less to the systemic circulation, thus reducing the danger of systemic side effects (LOE III; SOR C) (ROVC: 100%)

54. HIA chemotherapy should be practiced in highly specialized centers (LOE II, SOR B) (ROA: 100%)

55. HIA chemotherapy is administered through surgically placed ports or implantable continuous infusion pumps (LOE II, SOR B) (ROVC: 100%)

56. The most common chemotherapeutic scheme in use is floxuridine with dexamethasone. The latter agent is given to reduce the risk of toxic injury to the biliary tract (LOE II, SOR B) (ROVC: 94%)

57. Side effects of HIA chemotherapy with floxuridine and dexamethasone are intestinal mucosa ulceration and diarrhea, biliary toxicity, and hepatic parenchyma toxicity (LOE II, SOR B) (ROVC: 100%)

58. HIA chemotherapy is usually combined with systemic chemotherapy (LOE II, SOR B) (ROVC: 96%)

59. Evidence on the use of other chemotherapeutic regimens (5-FU with leucovorin, or oxaliplatin) is limited (LOE IV; SOR D) (ROVC: 95%)

60. Indications of HIA chemotherapy combined with systemic chemotherapy are: 1) failure of first-line systemic chemotherapy to convert unresectable liver metastases to resectable; 2) as a first-line approach for initially unresectable disease; and 3) as adjuvant approach after R0 resection of liver metastases (LOE IV; SOR D) (ROVC: 90%)

61. Although HIA chemotherapy may offer a longer PFS, the approach does not improve OS (LOE II; SOR B) (ROVC: 98%)

**Surgery**

**Background**

Surgery can be performed safely after 4 weeks from the last cycle of induction chemotherapy with or without cetuximab, and 5-8 weeks following chemotherapy with bevacizumab. A reasonable time interval allows liver function to recover [70,115]. Resection of the metastases should be performed as soon as metastases become resectable, since unnecessary prolonged administration of chemotherapy may lead to a higher postoperative morbidity. A R0 resection should be attempted including all initial sites of disease if possible, even in the case of complete clinical response, because “expectant policy” and deferral of surgery is associated with increased recurrence rates [116,117]. In order to preserve functional hepatic parenchyma, resection, either in the form of typical lobectomy or of multiple segmental and wedge resections, lesions <3 cm can be thermally ablated with the use of RFA or MWA, and with low added morbidity and mortality. Ablation of lesions >3 cm is associated with increased recurrence rates [56,118-123]. MWA carries the advantages of less risk of charring, less incidence of “heat sink” effect and ability to ablate larger lesions as compared to RFA. The disadvantage of MWA is the increased risk of thermal injury to adjacent structures [122,124,125].

In case of increased but resectable metastatic burden and increased risk of liver failure after one stage resection, a two-stage surgery can be applied (first stage: wedge resections of the lobe to remain and PVL of the branch of the lobe to be resected; second stage: hepatic resection). PVE and PVL show similar hypertrophy of the remnant hepatic lobe [126]. According to a meta-analysis including 1088 patients from 37 studies [127] and a more recent study [128], PVE is a safe and effective procedure that induces sufficient liver hypertrophy of the remnant liver and prevents postoperative liver failure.

A recently described alternative approach to induce rapid hypertrophy of the left lobe and enable extensive right lobectomy is the “in situ split” procedure combined with right PVE or PVL at a first stage and extensive right lobectomy at a second stage, at the same admission. There is little evidence on this approach, which is associated with mortality of 12% [129].

Portal vein chemotherapy at the immediate postoperative period, after R0 resection of liver metastases does not seem to be of any additional oncological benefit over curative surgery alone [130]. The postoperative morbidity is more related to the duration of the chemotherapy, than to the type of chemotherapy that is administered, although oxaliplatin and irinotecan may cause different histological changes in liver parenchyma, oxaliplatin is related to sinusoidal liver lesions [131-133].
Non-surgical management of liver metastatic disease

Non-surgical management of liver metastasis should not be pursued instead of resection of metastasis, but only attempted in case of unresectable liver disease [134,135]. Patients not suitable for surgery, due to either bulky lesions, or unfitness, or presence of non-resectable extrahepatic metastases, should be treated upfront with systemic chemotherapy. Patients showing progression of the disease after chemotherapy, or presenting toxicity from systemic therapy that limits chemotherapeutic options, are potential candidates for palliative, ablative or regional chemotherapeutic interventions.

Ablative techniques

**Background**

RFA. Patients whose tumors have been downsized by chemotherapy but remain still unresectable, can be subjected to ablative therapy. RFA is accepted as preferable to other ablative techniques for treating colorectal metastases. Factors determining success are lesion size, the number of lesions and location [136]. Ablation should be reserved for patients with a limited number of smaller tumors. Local recurrence is significantly higher for >3-5 colorectal metastases, while prognosis is worse when more than five metastases are present at the time of ablation [118,123]. Larger tumors may be treated with a combination of ablation and arterial embolization or chemo-embolization [137,138].

RFA may be performed intraoperatively, laparoscopically, or percutaneously. The percutaneous approach is associated with the least procedural risk and may be performed under local anesthesia, but is less effective. Percutaneous RFA should be considered as a primary treatment option for patients with unresectable hepatic tumors or conditions that prohibit general anesthesia or abdominal surgery [139]. Laparoscopic RFA has been also attempted with low local failure rates, a median recurrence free survival of 15 months, and mean actuarial survival of approximately 4 years [140].

**Other ablative modalities.** Other palliative modalities of liver metastatic disease are: 1) **external radiation.** Use of proton-based instead of stereotactic body radiotherapy for solitary unresectable liver metastatic lesions spares considerably liver parenchyma and prevents injury to adjacent structures [141-145]; 2) **radiochemoembolization.** Combination of chemoembolization (mitomycin, cisplatin and doxorubicin) with radioembolization induces a short-term reduction in tumor size [146], and 3) **portal vein chemotherapy.** These modalities are in use in highly specialized centers and applied in selected patients. At present, evidence is limited.

**RECOMMENDATIONS**

65. **Indications for ablation techniques to control colorectal metastatic disease are poorly defined (ROVC: 90%)**

66. **Ablation techniques can be applied as an adjunct to hepatic resection, to preserve liver parenchyma (LOE II; SOR B) (ROVC: 97%)**

67. **Ablation techniques can be recommended as a first-line treatment for small metastatic lesions at appropriate locations (not in the vicinity of large vessels), if surgical resection is not considered safe or contraindicated (LOE II; SOR B) (ROVC: 99%)**

68. **There is poor evidence to support ablation of liver metastases as a first-line treatment instead of surgical resection (LOE III, SOR D) (ROVC: 97%)**

69. **Application of ablation therapies are limited by the: 1) features of the lesions (size, location, multiplicity); 2) substantial morbidity; and iii) high recurrence rates (LOE II; SOR A) (ROVC: 97%)**

70. **Factors affecting recurrence after RFA are tumors >3 cm, tumor proximity to large vessels and percutaneous instead of surgical approach (LOE II; SOR B) (ROVC: 96%)**

Selective internal radiation therapy (SIRT)

**Background**

SIRT with Yttrium-90 of liver metastases from primary CRC often results in a high rate of tumor regression. All patients indicated for a SIRT must not have widespread metastases, which are ruled out by PET-CT or whole body CT/MRI. Current evidence on the safety of SIRT for non-resectable colorectal metastases in the liver is adequate. For patients who have previously been treated with chemotherapy, there is evidence that SIRT can achieve control of hepatic metastases, but the evidence of its effects on survival and on quality of life is inadequate (NICE consultation document 2011).

The response rate based only on changes in tumor volume according to the CT (RECIST) is often poor. However, after SIRT there is a significant decrease of tumor markers and FDG-uptake, proving that tumor nodules are no longer viable. Patient selection is also a key issue because there are considerations that a subgroup of patients with huge metastases or pre-existing extrahepatic manifestations seem to benefit less from this therapeutic modality [147]. There is no evidence that the modality can be applied as a first-line treatment in combination with systemic chemotherapy for unresectable liver metastases [148-151].
Prognostic factors of synchronous metastatic disease are metastatic disease share similar prognosis [25,160-163]. Supporting the view that synchronous and metachronous metastatic disease [28,155-159], although there are reports be associated with worse prognosis compared to metachronous

**Background**

**Synchronous primary and metastatic disease**

**General considerations**

Background

Synchronous primary CRC and metastatic disease seems to be associated with worse prognosis compared to metachronous metastatic disease [28,155-159], although there are reports supporting the view that synchronous and metachronous metastatic disease share similar prognosis [25,160-163]. Prognostic factors of synchronous metastatic disease are primary tumor differentiation and lymph node status, presence of complications from the primary lesions, number and size of metastatic lesions, potential of curative surgery of both primary and metastatic disease, presence of extrahepatic metastatic disease and the level of serum CEA [164].

After MDT discussion and continuous at every stage-interval assessment, as well as personalization of treatment, planning, timing and sequencing of therapeutic modalities, namely chemotherapy and intervention for primary and secondary disease, are critical for the best oncological outcome. The best outcomes are achieved after R0 resection of both primary and metastatic disease. Resection of the primary lesion leaving intact metastatic disease, does not improve oncologic outcome. In the presence of unresectable metastatic disease, primary lesion is resected only for cure of complications [165-168].

**Transarterial chemoembolization (TACE)**

**Background**

TACE using irinotecan-loaded beads is safe and effective in the treatment of patients following failure of standard systemic chemotherapy. Patients may receive repeat embolizations (max 100 mg per session). The technique is associated with minimal complication rate and acceptable tumor response [152,153]. Neither the number of liver lesions, the size of liver lesions or the extent of liver replacement (<25% vs. >25%) are predictors of OS. TACE may also achieve downsizing of metastases, so that they can be treated by thermal ablation, at a following stage [154]. Treatment by TACE shows a significant benefit for patients who have failed first- and second-line systemic chemotherapy, and is potentially a more effective approach when compared to the historical response rates after third- or fourth-line systemic chemotherapy. The fact that substantial extrahepatic progression is often observed after regional treatment for liver metastases, further suggests that systemic chemotherapy should be added to chemoembolization.

**Upfront systemic chemotherapy**

Upfront systematic chemotherapy is indicated in all cases. The most common schemes in use are FOLFOX and FOLFIRI [169]. The addition of targeting agents (i.e. bevacizumab) increases the risk of bleeding or perforation at the site of primary lesion [170]. Response to upfront chemotherapy is a prognostic marker for outcomes and a tool of patient’s selection for further treatment, preventing unnecessary non-therapeutic operations, in case of lack of response. Most hepatic metastatic lesions reduce in size or remain stable, and this translates to reduced recurrence rates, and better survival rates [33,169,171-174]. In general, upfront chemotherapy is considered safe for the intact primary lesion, unless complications are pending or present [175-177]. Also, upfront chemotherapy and delay of resection of rectal cancer does not impair oncological outcomes and might also prevent local recurrence [178]. Given the fact that mostly primary but also metastatic lesions may disappear as a result of favorable response to chemotherapy, it is recommended that the site of the lesions to be marked (Indian ink for the primary; metallic coils for the metastatic lesions) prior to treatment initiation [179].
Surgical approaches

There are three approaches to the sequence of surgical procedures: 1) simultaneous resection of both primary and metastatic disease; 2) surgery for the primary lesion first; and 3) surgery for the metastatic (hepatic) lesions. Sequencing of procedures is personalized. Priority is given to the most pressing component of the disease, attempting to minimization of morbidity and mortality rate.

Simultaneous resection of both primary lesion and metastatic disease carries less morbidity, similar mortality and less length of hospital stay as compared to staged approach, provided that colorectal surgery is straightforward and hepatic resection is limited to segmental (up to four segments) or wedge excisions [175,180-186]. When a major hepatectomy is required, simultaneous surgery is associated with worse short-term outcomes [187,188]. When major surgery for the primary lesion (i.e. low anterior resection of rectum, abdominoperineal resection of rectum, subtotal colectomy) or the hepatic metastatic disease (i.e. right lobectomy, extended liver resection), a staged approach is recommended [184]. It has been suggested that when the duration of simultaneous resection is anticipated to be <6 h and expected morbidity is acceptable, a simultaneous resection must be chosen. Otherwise, a staged approach is recommended [189].

At the staged approach either colorectal or hepatic surgery is performed at the first stage. Colorectal surgery “first” is chosen in case of perceived complications at the site of the primary lesion (perforation, bleeding obstruction), followed by systemic chemotherapy, followed by resection of liver metastases [180,185,186,188,190]. Liver surgery “first” is the approach of choice, when progression of metastatic disease to become unresectable is imminent. After systemic chemotherapy and resection of metastatic disease, chemotherapy, alone or in combination with radiotherapy in case of rectal cancer, is given [191]. Resection of the primary lesion follows [175,192,193].

In selected cases and in experienced centers, a laparoscopic approach can be applied both for the resection of primary and hepatic metastases, either by the one- or two-stage approach [194].

Location of the primary lesion

The approach of synchronous primary and metastatic colorectal disease is also personalized according to the location of the primary tumor, as follows:

Synchronous metastatic colon cancer (Fig. 3)

Initially resectable synchronous disease should be treated preoperatively with FOLFOX for 3 months, then by resection of the primary tumor and of metastases, either staged or synchronous, followed by postoperative administration of FOLFOX for 3 months.

a) Symptomatic primary disease: In case of symptomatic primary and resectable systemic disease, resection of the primary tumor, and metastases if possible, should be performed upfront, followed by “adjuvant” chemotherapy for 6 months. In case of symptomatic primary and unresectable systemic disease, resection of the primary tumor should be performed, followed by induction chemotherapy and further treatment according to response.

b) Asymptomatic primary disease: In case of unresectable systemic disease, there is no benefit from upfront surgery for a non-symptomatic primary tumor. Instead, upfront systemic chemotherapy is recommended. In case of secondary resectability of metastatic disease after induction chemotherapy, resection of the primary tumor and metastases either staged or synchronous, should be performed. In case of insufficient response of liver metastases to induction chemotherapy, surgical resection may not be performed, as tumor R0 resection does not improve survival.

Synchronous rectal tumor and metastatic disease (Fig. 4)

In case of a rectal tumor with synchronous metastatic disease, treatment should be personalized. There are the following options:

a) Non-symptomatic early or intermediate rectal tumor: chemotherapy followed by assessment of the response of the metastatic lesion. If response is favorable and resection of metastatic disease is feasible, surgery should be considered: 1) primary tumor surgery first, followed by metastatic disease resection at a second stage; 2) metastatic disease resection first, followed by resection of primary disease at a second stage; 3) one stage resection of primary and metastatic disease. If response of metastatic disease is not favorable and primary tumor is asymptomatic, no surgery is required.

b) Symptomatic advanced rectal tumor: Chemoradiotherapy followed by chemotherapy at the resting period and assessment of the response of both the primary and the metastatic lesion. Depending on the response, the following therapeutic measures are taken: If response of both primary and metastatic disease is favorable, with anticipated R0 resection of both: 1) resection of primary and metastatic disease at one stage; 2) metastatic disease resection first, followed by resection of primary disease at a second stage; 3) primary tumor surgery first, followed by metastatic disease resection at a second stage. If response of metastatic disease is not favorable and primary tumor becomes asymptomatic, no surgery is required. Also, if R0 resection of the primary lesion is not feasible no surgery is required. In both latter cases, palliative measures can be undertaken. If response of metastatic disease is not favorable and primary tumor is asymptomatic no surgery is required. Also, if R0 resection of the primary lesion is not feasible, no surgery is required. In both latter cases, palliative measures can be undertaken.
Prognosis of metastatic liver disease

3- and 5-year OS rates are around 50% and 30-47% respectively, after curative treatment of hepatic metastatic disease (R0 resection), while median survival extends to 50 months. Predictors of adverse oncological outcomes are female gender, primary disease with advanced T and N stage, number of hepatic metastatic lesions >4, size of largest lesion >5 cm, bilobar disease, extrapleural disease, serum CEA >5 ng/mL and serum CA19-9 >37 U/mL. Presence of positive lymph nodes at the hepatic hilum has an adverse effect on oncological outcomes [38]. Also, presence of C-reactive protein polymorphism and KRAS/BRAF mutations are adverse independent prognostic factors in patients with synchronous primary and metastatic disease [203].

It has been shown that a limited resection margin is associated with an increased risk of recurrence [204], and recurrence after attempted curative resection is attributed to an involved resection margin [205]. However, neither the extent of the resection margin nor the type of hepatectomy (anatomic or non-anatomic) is a significant prognostic factor for OS and DFS or local recurrence [200]. In addition, intraoperative increased blood loss, increased transfusion requirements and increased number of fresh frozen plasma transfusion, adversely affect immediate postoperative morbidity and mortality, as well as long-term oncological outcomes [201,206]. Furthermore, increased immediate postoperative morbidity seems to be associated with impaired OS [49,207]. Additional prognostic factors of impaired oncological outcomes are: 1) an increased systemic
inflammatory response [206]; 2) neutrophils/lymphocytes ratio >5 [208]; 3) a less than twice increase in monocytes count [197] and increased serum VEGF and TIMP-1 concentrations.

Patients after curative resection of hepatic metastases should be followed-up regularly for local recurrence (liver US, CT, MRI) and metachronous extra-hepatic lesions (CT, PET-CT). The rate of metachronous extra-hepatic metastases is 19% [209]. In case of recurrent liver metastases, repeat curative resection is associated with acceptable morbidity and low mortality rates. 5-year survival rate after repeat liver resection could be more than 45%, provided that a R0 resection has been achieved and there is no extra-hepatic disease [210,211]. Liver transplantation is rarely indicated (treatment related morbidity), and only in case of absent extra-hepatic disease [212].

### Table 3 Completeness of cytoreduction (CC) scoring

| Residual peritoneal disease | CC scoring |
|----------------------------|------------|
| No obvious disease         | CC=0       |
| Residual lesion ≤0.25 cm   | CC=1       |
| Residual lesion 0.25 - 2.5 cm | CC=2   |
| Residual lesion >2.5 cm    | CC=3       |

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*Figure 4* Strategy and treatment algorithm of synchronous rectal cancer and liver ± lung metastases

*CRM, circumferential resection margin; EMVI, extra-mural venous invasion; MDT, multi-disciplinary team; CRTx, chemo-radiotherapy*
RECOMMENDATION
Initially resectable synchronous disease

79. Preoperative chemotherapy helps select candidates for resections according to response. In most cases metastatic burden decreases or remains stable, and this response predicts better survival, reduced recurrence rate, and less non-therapeutic operations (LOE I, SOR A) (ROVC: 98%)

80. Initially resectable synchronous disease, should be treated preoperatively with FOLFOX or FOLFIRI for 3 months, followed by resection of the primary tumor and metastases, either staged or synchronous, and postoperatively by chemotherapy with FOLFOX or FOLFIRI for 3-months (LOE II; SOR B) (ROVC: 85%)

81. Preoperative chemotherapy should be limited (4-6 cycles), to reduce risk of potential hepatotoxicity (LOE II, SOR A) (ROVC 100%)

82. Preoperative chemotherapy, leaving primary tumor intact, is considered safe, unless complications are pending (LOE II, SOR A) (ROVC: 97%)

83. Prior to preoperative chemotherapy, marking of both primary lesion with Indian ink and hepatic metastases with metallic coils is recommended, as their location might be impossible after chemotherapy (LOE III, SOR D) (ROVC: 83%)

84. Bevacizumab, in preoperative chemotherapeutic regimes, increases the risk of bleeding and perforation, and should be avoided in ulcerated lesions (LOE I, SOR A) (ROVC: 88%)

85. Following induction chemotherapy, sequencing of procedures (simultaneous, or colorectal first, or liver first) should be personalized and based on MDT discussion. Priority should be given to the more pressing disease component (LOE III, SOR B) (ROVC: 99%)

86. Simultaneous resection of primary lesion and metastatic disease is recommended: 1) in case of a standard (not extended) colon resection with up to two segment hepatectomy; and 2) in case of extended colon or rectum resection with limited liver resection (LOE II, SOR B) (ROVC: 96%)

87. In case of symptomatic primary and resectable systemic disease, resection of the primary tumor, and metastases if possible, should be performed upfront, followed by “adjuvant” chemotherapy for 6 months (LOE IV; SOR C) (ROVC: 92%)

88. When a simultaneous resection is planned, liver resection is attempted first by colon or rectal resection (LOE II, SOR B) (ROVC: 82%)

89. Staged resection is suggested when major resections, both hepatic and colorectal, are anticipated (LOE II, SOR B) (ROVC: 100%)

90. Colon surgery is undertaken first in case of imminent complications (bleeding, obstruction, perforation). Following systemic chemotherapy,

RECOMMENDATION
Resection of metastatic disease is attempted (LOE II, SOR B) (ROVC: 98%)

91. In case of synchronous colon cancer and metastatic disease, resection of metastases is attempted first, when there is a risk of progression to an unresectable status if the patient remains off chemotherapy (LOE II, SOR B) (ROVC: 99%)

92. In case of synchronous rectal cancer and resectable metastatic disease of substantial burden, the following sequence is proposed: 1) systemic chemotherapy; 2) metastases resection; 3) chemotherapy or chemoradiation for the rectal tumor according to indication (personalization); 4) rectal tumor resection; and 5) systemic chemotherapy (LOE II, SOR B) (ROVC: 89%)

93. Simultaneous resection seems to be associated with similar mortality, less morbidity and shorter length of stay, compared to staged resection (LOE II, SOR B), although there is evidence that simultaneous resection could be associated with higher mortality (LOE II, SOR C) (ROVC: 97%)

RECOMMENDATIONS
Initially unresectable metastatic disease, becoming potentially resectable after chemotherapy

94. In case of initial unresectable and secondary resectable metastatic disease after induction chemotherapy, resection of the primary tumor and metastases, either staged or synchronous, should be performed (LOE IV; SOR D) (ROVC: 99%)

95. Higher rates of conversion of initially unresectable to resectable liver metastases are achieved, if systemic chemotherapy is combined with HIA, in highly specialized centers (LOE III, SOR C) (ROVC: 95%)

96. Following resection of eventually resectable metastases of rectal cancer, the primary rectal lesion should be treated according to local stage of the disease (resection of rectum or radiotherapy followed by resection of rectum) (LOE IV; SOR D) (ROVC: 99%)

97. In case of insufficient response of liver metastases to induction chemotherapy, surgical resection may not be performed, as primary tumor R0 resection does not improve survival (LOE III; SOR C) (ROVC: 87%)

RECOMMENDATIONS
Definitely unresectable metastatic disease

98. Systemic chemotherapy is recommended to prevent further progression of the metastatic disease and related symptoms (LOE IV; SOR D) (ROVC: 95%)
Metastatic CRC: surgical management guidelines

Synchronous liver and lung metastatic disease

After upfront systemic chemotherapy, R0 resection of both liver and lung metastases is associated with a reasonably high OS rate of more than 50% [213]. Survival also depends on the number of extra-hepatic metastatic deposits [72].

Management of peritoneal disease

Background

The incidence of peritoneal carcinomatosis in CRC is around 11% [214], and until recently, peritoneal carcinomatosis has been considered as a terminal stage of the disease, liable to palliative treatment only. Systemic chemotherapy monotherapy (5-FU) in peritoneal disease is associated with a median survival of 5.2-12.6 months [214-216].

Maximal cytoreductive surgery using standard peritonectomy procedures, combined with perioperative intraperitoneal chemotherapy, either as hyperthermic intraoperative (HIPEC) or as early postoperative under normothermia (EPIC), has demonstrated a significant improvement in survival rates, in selected patients. Median survival of around 19 months and 3- and 5-year OS of 39% and 19% respectively have been reported by several studies, most of them of low quality [216-228], and one meta-analysis [229].

Cytoreductive surgery with EPIC or HEPIC does not offer benefit to all patients with peritoneal carcinomatosis of colorectal origin, and proper patient selection is required. Extent of peritoneal involvement, as calculated by the Peritoneal Cancer Index (PCI) (Fig. 5) [216,217,219,221,223-226,230], and completeness of cytoreduction (CC), assessed by a specific CC-score, are the most important variables to determine oncological outcomes (Table 3). Bad prognostic factors are a PCI>13, the presence of peritoneal nodules >5 cm at the ligament of Treitz or adjacent to the root of jejunal mesentery and multiple segmental small bowel obstructions. Only, a R0 cytoreduction is of oncological benefit. Additional factors determining overall outcomes are good general PS, normal hematological profile, normal renal and liver function, absence of distant unresectable metastatic disease (liver, bone marrow, lung, brain), and absence of a second malignancy at risk for recurrence [216,217,219,221-227,231].

Cytoreductive surgery

Currently, peritonectomies include: 1) epigastric peritonectomy; 2) right and left subdiaphragmatic peritonectomy; 3) greater omentectomy and splenectomy; 4) lesser omentectomy; 5) cholecystectomy with resection of the omental bursa; 6) right and left lateral peritonectomy; 7) pelvic peritonectomy; and 8) resection of other organs if required to achieve a R0 resection.

Perioperative intraperitoneal chemotherapy

Even if the macroscopically visible tumor has been completely removed after maximal cytoreductive surgery, microscopic residual tumor at the peritoneal surfaces may be present, and secondary peritoneal tumors may appear within 2-3 years after initial surgery. The rationale of intraperitoneal chemotherapy is based on the fact that cytostatic agents exhibit a penetration of approximately 1-2 mm into tissues, and thus may result to the eradication of the microscopic residual tumor.

HIPEC enhances cytotoxicity and improves drug penetration. Heat itself has potential antitumor properties. HIPEC may be performed by an open or a closed technique with the use of specific devices. The most frequently cytotoxic drugs in use are mitomycin-C, cisplatin, doxorubicin, and oxaliplatin. Renal toxicity of intraperitoneal chemotherapy is avoided by careful monitoring of urine output during perfusion. EPIC under normothermia is used with the same intent as HIPEC, within the first five postoperative days after cytoreduction before intra-abdominal adhesion formation. Distribution and effectiveness of cytostatic drugs to penetrate the peritoneal-plasma barrier are inferior to that of HIPEC.

Hospital mortality after cytoreductive surgery and HIPEC or EPIC is 0-4%. Hospital morbidity is high, ranging from 20% to 54%, implying that the method must be performed at highly specialized centers [218,220,221,232,233]. According to

RECOMMENDATIONS

Definitely unresectable metastatic disease

99. If first-line treatment fails to control progression of metastatic disease, loco-regional treatment, in the form of HIPEC chemotherapy, chemoradiation, chemoembolization, or ablative modalities, can be used, in highly specialized centers (LOE IV; SOR D) (ROVC: 94%)

100. In case of a symptomatic primary lesion and synchronous unresectable systemic disease, resection of or other intervention (brachytherapy, stenting, etc) to the primary lesion should be performed, followed by systemic chemotherapy and further treatment according to response (LOE IV; SOR D) (ROVC: 96%)

101. In case of unresectable systemic disease, there is no benefit from upfront surgery for a nonsymptomatic primary tumor. Instead, upfront systemic chemotherapy is recommended (LOE IV; SOR D) (ROVC: 97%)
current literature, the 3-year survival rate ranges between 23% and 47%. Patients in whom complete cytoreductive surgery was possible, had a median survival far superior than that in patients with incomplete cytoreductive surgery. Systematic chemotherapy should always be given after cytoreduction and intraperitoneal chemotherapy to improve oncological outcomes [217,221,227,234-236].

It appears that cytoreductive surgery with HIPEC or IPEC has a survival benefit over palliation with systemic chemotherapy alone. However, cytoreduction combined with HIPEC or IPEC does not seem to offer any additional benefit over cytoreduction with systemic chemotherapy. Patients with resectable liver metastases should not be excluded from cytoreduction and HIPEC or IPEC [229,237].

**RECOMMENDATIONS**

102. In patients with peritoneal carcinomatosis of colorectal origin, the option of being treated with maximal cytoreductive surgery, combined with perioperative intraperitoneal chemotherapy, and followed by systemic chemotherapy, can be considered (LOE II; SOR B) (ROA: 97%)

103. The procedures should be performed in specific centers (LOE II; SOR B) (ROA: 99%)

104. Best oncological outcomes are obtained when a R0 cytoreduction is achieved and a limited extension of intraperitoneal disease is present, in fit patients (LOE II; SOR B) (ROA: 100%)

**Bone metastases**

CRC metastases only to bones are extremely rare, and are usually associated with lung metastases. Resistant lung metastases predict potential disease progression to the bones, more than liver metastases. Treatment is palliative and consists of systemic chemotherapy and localized treatment (radiation, cyber-knife) to treat symptoms [238].

**Follow up after curative resection of primary disease**

Patient follow up depends on stage of the disease, quality of surgery, and amenability for intervention with either resection of recurrent disease or consideration of further systemic therapy. Four recent meta-analyses have shown that intensive follow up improves OS by 7-13% and is now considered as standard [239]. Generally, valid assessments are 3-monthly clinical visits for the first three years, followed by 6-monthly visits for further two years with clinical examination, evaluation of long-term toxicities (polyneuropathy after oxaliplatin), and CEA testing [240].

Complete colonoscopy must be performed at initial diagnosis, after three and afterwards every five years. If the diagnostic colonoscopy was incomplete, this should be repeated at 6 months after surgery. If follow-up colonoscopy shows advanced adenoma, successfully and completely excised, it should be repeated in one year.

In patients with high-risk disease, CT scan of the chest and abdomen every 6-12 months could be considered, although such close follow up should be confined to patients possibly amenable to resection of hepatic or pulmonary recurrence. Follow up CT scans should be performed with the same imaging protocols and contrast phases of enhancement. If MRI was used for the initial staging, MRI should also be used for the follow up, because CT images cannot be compared to MRI images due to different sensitivity/specificity [241]. Liver contrast-enhanced US could substitute for abdominal CT scan regarding follow up of liver metastases, particularly in young patients with no evidence of extra-hepatic disease [239,242]. Finally, routine PET-CT scanning is not recommended for surveillance [243].
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