A Comprehensive Review of Clinical Trials on EGFR Inhibitors Such as Cetuximab and Panitumumab as Monotherapy and in Combination for Treatment of Metastatic Colorectal Cancer

Mohammad Hossein Yazdi 1, Mohammad Ali Faramarzi 2, Shekoufeh Nikfar 3, and Mohammad Abdollahi 4*

1. Department of Research and Development, Pasteur Institute of Iran, Karaj, Iran
2. Department of Pharmaceutical Biotechnology, Faculty of Pharmacy and Biotechnology Research Center, Tehran University of Medical Sciences, Tehran, Iran
3. Department of Pharmacoeconomics and Pharmaceutical Administration, Faculty of Pharmacy, Tehran University of Medical Sciences, Tehran, Iran
4. Department of Toxicology and Pharmacology, Faculty of Pharmacy and Pharmaceutical Sciences Research Center, and Endocrinology & Metabolism Research Center, Institute of Clinical Endocrine Sciences, Tehran University of Medical Sciences, Tehran, Iran

Abstract

Metastatic colorectal cancer is the fourth most common cause of death due to cancer after those of lung, stomach, and liver. Anti epidermal growth factor receptor drugs as a targeting therapy seem to be good candidates for curing metastatic colorectal cancer. Two available anti epidermal growth factor receptor monoclonal antibodies are cetuximab and panitumumab which have been approved for metastatic colorectal cancer treatment. Through the available literature on NCBI and clinical trials, 31 clinical trials in which cetuximab or panitumumab as monotherapy or in combination with chemotherapy were used for the treatment of metastatic colorectal cancer patients in different line settings and 12 clinical trials in which bevacizumab was used for being compared with anti epidermal growth factor receptor monoclonal antibodies or chemotherapy were chosen for reviewing and comparing the results of overall survival, progression free survival and adverse effects. Cetuximab and panitumumab are well accepted for the treatment of mCRC patients at all stages in different line settings. Although cetuximab administration in metastatic colorectal cancer patients is mostly associated with better overall survival and panitumumab results in better progression free survival, to confirm the superiority of each of them in the treatment protocol of epidermal growth factor receptor monoclonal antibodies, more clinical trials with larger sample size are needed. Through current available data from clinical studies, it can be concluded that the best treatment outcome is achieved by a combination of anti epidermal growth factor receptor monoclonal antibodies with conventional chemotherapy.

Keywords: Anti-EGFR drugs, Cetuximab, Metastatic colorectal cancer, Panitumumab

Introduction

Despite all advances in the therapeutic modalities for Colorectal Cancer (CRC), this malignancy continues to be the fourth most common cause of cancer death after lung, stomach, and liver cancer. Consequently, the overall five-year survival remains very poor about 10% for patients at metastatic stage of this disease (mCRC). However, with the introduction of several chemotherapeutic agents such as irinotecan and oxaliplatin in combination with fluorouracil or leucovorin, the hope for survival of mCRC patients has been created and development of drug resistance results in investigating other therapeutic options. The abnormal extra expression of the Epidermal Growth Factor Receptor (EGFR) is frequently associated with many human malignancies including mCRC. Therefore, anti-EGFR drugs seem proper candidates for the treatment of mCRC using targeted therapy. Currently, two clinically available anti-EGFR Monoclonal antibodies (Mabs) are cetuximab and panitumumab, which reached FDA approval in 2004 and 2007 respectively, for the treatment of mCRC. Cetuximab is a chimeric mouse/human Mab given by intravenous infusion, binds to the EGFR, stops the binding and activation of the downstream signaling pathways and prevents the...
cell proliferation, invasion, metastasis, and neovascularization (Box 1) ⁷. Panitumumab is a fully human Mab with the same properties as cetuximab (Box 2). Clinical use of anti-EGFR Mabs has shown variable data in different line settings of mCRC therapy. For instance, some studies have demonstrated a survival benefit of single-agent therapy of cetuximab or panitumumab in the third line setting ⁸,⁹ while another study showed the superiority and better overall survival of cetuximab-irinotecan combination in comparison to panitumumab monotherapy in previously treated mCRC patients ¹⁰. However, the heterogeneity of study population and many other factors may justify these various results.

In the present review, in order to better understand the advantage and/or disadvantage of application of anti-EGFR therapy in different line settings of mCRC treatment, both as monotherapy and in combination with chemotherapy, and corresponding adverse effects, recent clinical trials were compared with each other.

**Comparison of cetuximab and panitumumab in combination or monotherapy of mCRC**

Most of clinical trials include the anti-EGFR therapy in the treatment protocol of mCRC in conventional chemotherapy refractory patients. In this regard, anti-EGFR Mabs are usually used in combination with some chemotherapeutic agents in second or third line setting for treatment. However, in some clinical trials after chemotherapy failure, anti-EGFR Mabs have been used in monotherapy. An important factor to select the cetuximab or panitumumab for the treatment of mCRC patients is the Kirsten Rat Sarcoma viral oncogene (KRAS) mutation status. In fact, mutation in the KRAS gene is considered as a negative predictor of response to cetuximab and/or panitumumab. In a study conducted on Japanese patients with mCRC, the response rate (RR) to cetuximab plus irinotecan therapy was around 17.9 and 0% in the KRAS wild-type (WT KRAS) and mutant subgroups, respectively ¹¹. Studies about the application of panitumumab in mCRC patients also

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**Box 1. Drug summary**

<table>
<thead>
<tr>
<th>Drug name</th>
<th>Cetuximab</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phase</td>
<td>Launched</td>
</tr>
<tr>
<td>Indication</td>
<td>For treatment of EGFR-expressing metastatic colorectal cancer</td>
</tr>
<tr>
<td>Pharmacology description</td>
<td>Epidermal growth factor receptor binding FAB</td>
</tr>
<tr>
<td>Route of administration</td>
<td>Intravenous</td>
</tr>
<tr>
<td>Protein structure</td>
<td>Not available</td>
</tr>
</tbody>
</table>

Protein chemical formula: C₆₄₈₄H₁₀₀₄₂N₁₇₃₂O₂₀₂₃S₃₆

Pivotal trial(s): [11, 12, 15]

The protein structure and the chemical formula are adapted from [http://www.drugbank.ca/drugs/DB00002](http://www.drugbank.ca/drugs/DB00002)

**Box 2. Drug summary**

<table>
<thead>
<tr>
<th>Drug name</th>
<th>Panitumumab</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phase</td>
<td>III</td>
</tr>
<tr>
<td>Indication</td>
<td>For treatment of EGFR-expressing metastatic colorectal cancer</td>
</tr>
<tr>
<td>Pharmacology description</td>
<td>Panitumumab (ABX-EGF) is a recombinant human IgG2 monoclonal antibody that binds specifically to the human epidermal growth factor receptor (EGFR)</td>
</tr>
<tr>
<td>Route of administration</td>
<td>Intravenous</td>
</tr>
<tr>
<td>Protein structure</td>
<td>Not available</td>
</tr>
<tr>
<td>Protein chemical formula</td>
<td>C₆₃₉₈H₉₈₇₈N₁₆₉₄O₂₀₁₆S₄₈</td>
</tr>
<tr>
<td>Pivotal trial(s)</td>
<td>PRIME trial [18]</td>
</tr>
</tbody>
</table>

The protein chemical formula and description are adapted from [http://www.drugbank.ca/drugs/DB01269](http://www.drugbank.ca/drugs/DB01269)
demanded the same results. The first line treatment options for patients with mCRC are varied and range from single drug to highly effective four-drug combination, but the best option is designated based on tumor and patient related factors and differs case by case. Cetuximab can be involved in the first line treatment of mCRC patient.

In one study of 845 patients with KRAS wild-type tumors, adding cetuximab to the first line chemotherapy, significantly led to an improvement in Overall Survival (OS) compared with chemotherapy alone. Also in another study, higher Progression Free Survival (PFS) was observed in WT KRAS patients treated with cetuximab in the first line chemotherapy. In addition, the combination of cetuximab with irinotecan/5-fluorouracil/folinic acid (FOLFIRI) in the initial treatment of mCRC in another clinical trial was not only well tolerated but also in one-quarter of patients changed the status of metastases from unresectable to resectable. Although it should be noticed that the combination of anti-EGFR with chemotherapeutic agents is not always allowed to be prescribed and some combinations should be avoided (e.g., oral or bolus fluoropyrimidines, oxaliplatin and cetuximab or panitumumab), but many studies indicate the benefits of combination therapy in the treatment of mCRC in different line settings.

In a randomized controlled trial of cetuximab plus chemotherapy for patients with WT KRAS unresectable colorectal liver-limited metastases, the combination of cetuximab with FOLFIRI (fluorouracil, leucovorin, and oxaliplatin) in one arm and chemotherapy alone in another arm were compared and results showed that patients in the combination arm had improved objective response rates (57.1 vs. 29.4%; p<0.01), increased 3-year OS rate (41 vs. 18%; p=0.013) and prolonged median survival time MST (30.9 vs. 21.0 months; p=0.013). In this study, patients who had resection of liver metastases had a significantly improved MST (46.4 vs. 25.7 months; p<0.01) compared with those who did not undergo surgery. However, the combination of cetuximab with chemotherapy is not always associated with positive response; in contrast to above studies, the results of EPOC trial raised strong doubt about this strategy as in this experiment, patients with operable metastases from colorectal cancer were randomized to receive fluoropyrimidine and oxaliplatin with or without cetuximab for 12 weeks before and then 12 weeks following surgery. In patients with resectable liver metastases, progression free survival was significantly worse in the cetuximab plus arm [14.8 vs. 24.2 months, Hazard Ratio (HR) (95% Confidence Interval (CI)] 1.50037 (1.000707 to 2.249517) p<0.048]. Wherein most of the clinical trials used cetuximab in the first line setting of their treatment protocol, cetuximab in the randomized clinical trial also resulted in a significant improvement in overall survival from 4.8 months to 9.5 months.

Application of panitumumab in the first line treatment of mCRC in combination with chemotherapy also created the hope for a better treatment outcome. In a PRIME phase III trial, to compare the combination of panitumumab with folinic acid, 5 fluorouracil and oxaliplatin (FOLFOX4) and FOLFOX4 alone in mCRC patients, in the first line setting, panitumumab has demonstrated significant improvement in progression free survival (median PFS, 9.6 vs. 8.0 months, respectively; HR, 0.80; 95% CI, 0.66 to 0.97; p=0.02), though the overall survival has not increased significantly in panitumumab-FOLFOX4 versus FOLFOX4 (median OS, 23.9 vs. 19.7 months, respectively; HR, 0.83; 95% CI, 0.67 to 1.02; p=0.072). Randomized phase III study of panitumumab with fluorouracil, leucovorin, and irinotecan (FOLFIRI) compared with FOLFIRI alone as second line treatment in patients with metastatic colorectal cancer also showed only improvement in PFS of patients with combination therapy (HR=0.73; 95% CI, 0.59 to 0.90; p=0.004); median PFS was 5.9 months for panitumumab-FOLFIRI versus 3.9 months for FOLFIRI, overall survival has not been changed significantly and median OS was 14.5 months versus 12.5 months, respectively (HR=0.85, 95% CI, 0.70 to 1.04; p=0.12). Administration of panitumumab in monotherapy regiment for WT KRAS mCRC patients following cetuximab-based regimens resulted in 67% disease control rate and 30% objective response rate, with meaningful change in PFS (4.2 months) and OS (9.6 months). The combination of panitumumab with decitabine (a hypomethylating agent) was also well tolerated and showed activity in previously cetuximab treated mCRC patients. Some studies also compared the application of panitumumab with bevacizumab (anti VEGF monoclonal antibody) for the treatment of mCRC. In a phase II PEAK study, to compare the FOLFOX regime in combination with either panitumumab or bevacizumab in 285 previously untreated mCRC patients (first line treatment), results indicated the similar Overall Response Rate (ORR). The PFS was also similar between arms.

In a study to compare the panitumumab monotherapy with cetuximab and irinotecan combination therapy as third line treatment setting in patients with KRAS wild-type mCRC, median overall survival was 7.7 months for the panitumumab group and 8.3 months in the cetuximab-irinotecan group and the survival outcomes were similar regardless of the therapy selected (HR:1.28; p=0.34). In ASPECTT trial, to compare cetuximab and panitumumab in mCRC chemorefractory patients (as monotherapy in third line setting), the application of cetuximab resulted in a bit lower overall response rate than panitumumab (ORR: cetuximab 19.8% and panitumumab 22%), while, the Progression Free Survival (PFS) of cetuximab was around 3 months longer (PFS: 4.4 vs. 4.1 months. HR: 1.00, 95% CI: 0.88-1.14). The overall survival of both drugs was equal (OS: cetuximab10.0 and panitumumab 10.4).
months. HR:0.97, 95% CI:0.84-1.11, p=0.0007). Considering achieved results, non-inferiority endpoint was met in ASPECT trial 22.

Although several studies are available to compare the effectiveness of cetuximab and panitumumab in mCRC patients (Table 1), regarding these mentioned trials, it can be concluded that cetuximab in combination with chemotherapy can offer better overall survival when compared to panitumumab plus chemotherapy. On the other side, in many randomized clinical trials, significant improvements in tumor response rates and progression free survival have been observed when panitumumab is combined with chemotherapy or used as monotherapy in chemorefractory mCRC individuals. Therefore, the priority of cetuximab or panitumumab and the decision of treatment option between these two available anti-EGFR Mabs depend mostly on the patient condition and clinical availability which should be considered by physicians. Regarding the role of KRAS status in the response rate of anti-EGFR therapy, it is fundamentally important to have an improved patient selection through the use of novel predictive biomarkers.

Comparison of the side effects of cetuximab and panitumumab in combination or monotherapy of mCRC

In contrast to traditional chemotherapeutic drugs, anti-EGFR Mabs do not cause systemic toxicities including nausea, vomiting, diarrhea, alopecia, and bone marrow suppression. However, their most common adverse effects are skin rash 23, hypomagnesemia 24, hypokalemia 25 and infusion reaction 26. Different grades of these side effects can be observed among mCRC patients treated with these drugs. Severe infections (≥grade 3) were sometimes reported in cetuximab and/or panitumumab trials as a treatment side effect, which caused the disruption of treatment, and even led to

### Table 1. The clinical outcome of cetuximab and panitumumab administration as monotherapy and in combination with chemotherapy in mCRC patients

<table>
<thead>
<tr>
<th>Clinical Trial</th>
<th>Treatment option</th>
<th>Patients</th>
<th>Endpoint</th>
<th>p-value</th>
<th>Ref</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kennecke H et al 2013</td>
<td>Mono PMab</td>
<td>178 (141 PMab, 37 CMab Combo)</td>
<td>OS: 7.7 vs. 8.3 Months</td>
<td>0.03</td>
<td>10</td>
</tr>
<tr>
<td>Price et al 2014 ASPECT</td>
<td>Mono CMab vs. Mono PMab in refractory mCRC</td>
<td>999 (499 PMab and 500 CMab)</td>
<td>OS: 10.0 vs. 10.4 Months PFS: 4.4 vs. 4.1 Months ORR: 19.8% vs. 22%</td>
<td>0.0007</td>
<td>12</td>
</tr>
<tr>
<td>Bokemeyer C et al 2012 CRYSTAL and OPUS</td>
<td>CMab Combo</td>
<td>845 KRAS wild-type tumors</td>
<td>OS: Improved PFS: Improved ORR: Improved</td>
<td>0.0062</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Pietrantonio F et al 2013</td>
<td>Mono PMab</td>
<td>30 Camab treated KRAS WT mCRC</td>
<td>DCR: 67% ORR: 30% PFS: 4.2 Months OS: 9.6 Months</td>
<td>0.12</td>
<td>19</td>
</tr>
<tr>
<td>NCT00115765 2013</td>
<td>BMab Combo PMab with BMab Combo</td>
<td>BMab+Oxaliplatin 410 PMab with BMab+Oxaliplatin 413</td>
<td>OS: BMab Combo=24.5 Months and PMab with BMab Combo=19.4 PFS: BMab Combo=11.4 Months and PMab with BMab Combo=10 Months</td>
<td>0.005</td>
<td>34</td>
</tr>
<tr>
<td>PEAK trial 2014</td>
<td>PMab Combo BMab Combo</td>
<td>278</td>
<td>OS: PMab Combo=34.2 Months BMab Combo=24.3 Months PFS: Similar between two arms</td>
<td>0.009</td>
<td>47</td>
</tr>
<tr>
<td>Santos-Ramos B et al 2013</td>
<td>Cmab Combo BMab Combo</td>
<td>227</td>
<td>PFS: CMab Combo=11.7 Months BMab Combo=9.6 Months</td>
<td>-</td>
<td>48</td>
</tr>
<tr>
<td>Pietrantonio F et al 2012</td>
<td>CMab Combo BMab Combo</td>
<td>96</td>
<td>OS: CMab Combo=22.7 Months BMab Combo=18.7 Months</td>
<td>0.55</td>
<td>50</td>
</tr>
<tr>
<td>Modest DP et al 2012</td>
<td>CMab Combo BMab Combo</td>
<td>54 KRAS p.G13D mutated patients</td>
<td>ORR: CMab=58% BMab=57%</td>
<td>-</td>
<td>51</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>PFS: CMab=8.0 Months BMab=8.7 Months OS: CMab=20. Months BMab=14.9 Months</td>
<td>0.9</td>
<td></td>
</tr>
<tr>
<td>Dotan E et al 2012</td>
<td>CMab+BMab Combo, CMab Combo</td>
<td>23: 12 CMab+BMab Combo, And 11 Cmab Combo</td>
<td>ORR: CMab+BMab Combo=36% CMab Combo=72%</td>
<td>0.72</td>
<td>52</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>PFS: CMab+BMab Combo=8.7 Months CMab Combo=14.4 Months OS: CMab+BMab Combo=18 Months CMab Combo=42.5 Months</td>
<td>0.52</td>
<td></td>
</tr>
<tr>
<td>EPOC trial 2014</td>
<td>Chemotherapy CMab Combo</td>
<td>257 patient with resectable colorectal liver metastases</td>
<td>PFS: Chemotherapy=14.1 Months CMab Combo=20.5 Months</td>
<td>0.003</td>
<td>58</td>
</tr>
</tbody>
</table>
pulmonary embolism and fatalities. However, many of these side effects also seem to be related to chemotherapeutic part of treatment regimen when combination of anti-EGFR with routine chemotherapy is considered. In this section of study, some reported side effects of cetuximab and panitumumab clinical trials were reviewed and these events between different arms of treatment in these studies were compared (Table 2).

In Amgen trials, to study the panitumumab monotherapy in mCRC patients with a history of treatment with fluoropyrimidine, irinotecan, and oxaliplatin chemotherapy from the 203 enrolled patients, who received intravenous (IV) infusion of panitumumab at a dose of 6 mg/kg once every 2 weeks, most reported adverse effects were abdominal pain, dehydration and dyspnea. In a randomized, multicenter, open-label, non-inferiority phase 3 study conducted by Price et al, in order to compare the efficacy of cetuximab and panitumumab in chemotherapy refractory mCRC patients when drug schedule was panitumumab for 499 patients (6 mg/kg once every 2 weeks) in

<table>
<thead>
<tr>
<th>Study</th>
<th>Treatment</th>
<th>Patients</th>
<th>Most common side effect</th>
<th>Ref</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cao Y et al 2010</td>
<td>Mono Cmab</td>
<td>3081</td>
<td>Grade 3 and 4 Hypomagnesemia (5.6%; 95% CI=3.0-10.2). All grade Hypomagnesemia (36.7%; 95% CI=22-54.4).</td>
<td>24</td>
</tr>
<tr>
<td>Cao Y et al 2010</td>
<td>Mono Cmab</td>
<td>1324</td>
<td>Grade 3 and 4 Hypokalemia [6.2% (95% CI 4.9-7.7)]. All-grade Hypokalemia [8.0% (95% CI 4.5-13.9)]</td>
<td>25</td>
</tr>
<tr>
<td>Amgen trial 2013 NCT00089635</td>
<td>Mono Pmab</td>
<td>203</td>
<td>Abdominal pain, Dehydration and Dyspnea 6/203 (2.96%)</td>
<td>29</td>
</tr>
<tr>
<td>Merck KGaA CRYSTAL trial 2014</td>
<td>CMab Combo</td>
<td>600</td>
<td>Diarrhea 36/600 (6.00%) Catheter related infection 8/600 (1.33%) Central line infection 10/600 (1.67%) Hypomagnesaemia 13/600 (2.17%)</td>
<td>36</td>
</tr>
<tr>
<td>NCT00083720 2011</td>
<td>Mono Cmab</td>
<td>85</td>
<td>Death 1/85 (1.18%). Dyspnea 3/85 (3.53%)</td>
<td>37</td>
</tr>
<tr>
<td>Herbert Hurwitz clinical trial 2013 NCT00290615</td>
<td>CMab+Bmab Combo</td>
<td>30</td>
<td>Diarrhea 6/30 (20.00%)</td>
<td>38</td>
</tr>
<tr>
<td>NCT00252564 2011</td>
<td>CMab+Bmab Combo</td>
<td>1324</td>
<td>Anemia in CMab 28/121 (23.14%) and in Bmab 45/118 (38.14%) Leucopenia in CMab 26/121 (21.49%) and in Bmab 37/118 (31.36%) Neutropenia in CMab 19/121 (15.70%) and in Bmab 49/118 (41.53%) Thrombocytopeny in CMab 22/121 (18.18%) and in Bmab 42/118 (35.59%)</td>
<td>39</td>
</tr>
<tr>
<td>OPUS clinical trial 2011 NCT000125034</td>
<td>CMab Combo</td>
<td>170</td>
<td>Pulmonary embolism 5/170 (2.94%) Anorexia 3/170 (1.76%)</td>
<td>40</td>
</tr>
<tr>
<td>NCT00193219 2013</td>
<td>CMab+Bmab Combo</td>
<td>31</td>
<td>Thrombosis/Thrombus/Embolism 4/31 (12.90%)</td>
<td>41</td>
</tr>
<tr>
<td>NCT00115764 2011</td>
<td>CMab+Bmab Combo</td>
<td>518</td>
<td>Fibril neutropenia 20/518 (3.86%) Nausea 24/518 (4.63%) Vomiting 27/518 (5.21%) Sepsis 16/518 (3.09%)</td>
<td>42</td>
</tr>
<tr>
<td>M.D. Anderson Cancer Center NCT00354978 2011</td>
<td>BMab Combo</td>
<td>43</td>
<td>Alopecia 22/43 (51.16%) Nausea, Fatigue and Hemorrhage 19/43 (44.19%)</td>
<td>44</td>
</tr>
<tr>
<td>Yamaguchi K et al 2014</td>
<td>Mono Cmab</td>
<td>2126</td>
<td>Infusion reactions in114 patients (5.7%). Grade 3-4 Infusion reactions in 22 patients (1.1%).</td>
<td>56</td>
</tr>
<tr>
<td>PETACC-8 2014</td>
<td>Mono Cmab CMab Combo</td>
<td>791</td>
<td>Grade 3 or 4 Acne-like rash: CMab=27% CMab Combo=≤1% Diarrhea: 14% vs. 9% Mucositis: 63% vs. 10% Infusion-related reactions: 55% vs. 30%</td>
<td>57</td>
</tr>
<tr>
<td>EPOC trial 2014</td>
<td>Chemotherapy CMab Combo</td>
<td>257</td>
<td>Pulmonary embolism and Death: Chemotherapy =1 case, CMab Combo=3 cases</td>
<td>58</td>
</tr>
<tr>
<td>COIN-B trial 2014</td>
<td>Intermittent Mono CMab Continuous Mono CMab</td>
<td>64 patients Intermittent Mono CMab 22% Continuous Mono CMab=29%</td>
<td>Grade 3, 4 Skin rash: Chemotherapy=15%, CMab Combo=15%</td>
<td>59</td>
</tr>
</tbody>
</table>

Most common adverse effects of cetuximab and panitumumab therapy as a single agent or in combination with chemotherapy in recent clinical trials. PMab: Panitumumab, CMab: Cetuximab, BMab: Bevacizumab, FOLFOX: Oxaliplatin, 5-fluorouracil and leucovorin, FOLFIRI: Fluorouracil, leucovorin and Irinotecan, Combo: Combination with Chemotherapy. Mono: Monotherapy of Mab
one arm and cetuximab for 500 patients (initial dose 400 mg/m²; 250 mg/m² once a week thereafter) in another arm, the incidence of adverse events of any grade and grade 3-4 was equal between two arms. Grade 3-4 skin toxicity occurred in 62/499 (13%) patients in panitumumab treated group and 48/500 (10%) patients in cetuximab group. The chance of grade 3-4 infusion reactions was lesser in patients treated by panitumumab than cetuximab ([1% < 0.5%] patient vs. [9% (2%) patients]) and the occurrence of grade 3-4 hypomagnesemia was higher in the panitumumab group ([35 (7%) vs. 13 (3%)]). Also one treatment-related fatal adverse event, a lung infection in a patient given cetuximab was observed. In a post-marketing surveillance study of panitumumab monotherapy in 3085 Japanese patients with mCRC, the most common adverse drug reaction was skin disorders (78.4%). Also, the chance of all grades of electrolyte abnormalities was 19.3%, interstitial lung disease was 1.3% (mortality chance of 0.6%), infusion reaction was 1.5%, and cardiac disorders was about 0.2% vs. 0.00%. A noticeable fact which has been observed in different clinical trials to study the effect of anti-EGFR antibodies in mCRC treatment is that patients with grade 2 or 3 skin-related adverse events had higher response rates and longer PFS than patients with grade 1 events. These findings are consistent with studies associating skin toxicity with response to anti-EGFR antibodies. Therefore, the selection of treatment protocol among available options should be done by considering the patient’s condition and probable chance of treatment success.

The addition of panitumumab to bevacizumab plus irinotecan or oxaliplatin based chemotherapy for mCRC patients also resulted in an increased risk of febrile neutropenia (Pmab. plus Bmab with chemotherapy as first arm: 20/518 (3.86%) and Bmab with chemotherapy as second arm: 9/510 (1.76%), diarrhea ([63/518 (12.16%) vs. 15/510 (2.94%)], nausea ([24/518 (4.63%) vs. 7/510 (1.37%)] and vomiting ([27/518 (5.21%) vs. 10/510 (1.96%)]). In another experiment, to study the effect of chemotherapy and bevacizumab with or without panitumumab in the first line treatment of mCRC, results showed that those individuals who received panitumumab in combination with bevacizumab and chemotherapy experienced a higher incidence of death (9% vs. 4%) vs. 4.63%)

Cetuximab in several clinical trials also showed different adverse effect. In CRYSTAL clinical trial with 600 mCRC patients, to compare the efficacy of cetuximab plus FOLFIRI and FOLFIRI chemotherapy alone, the most common observed side effects were as follows: diarrhea ([36/600 (6.00%) in Cmb vs. 21/602 (3.49%) in FOLFIRI], catheter related infection [8/600 (1.33%)] vs. 1/602 (0.17%), central line infection ([10/600 (1.67%) vs. 4/602 (0.66%)] and hypomagnesemia ([13/600 (2.17%) vs. 1/602 (0.17%)]). In another study using cetuximab for the treatment of 85 mCRC patients, one case of treatment related death was observed and the rate of dyspnea was around 3/85 (3.53%). Also, cetuximab in combination with capcitabine, oxaliplatin, and bevacizumab for the treatment of 30 mCRC patients caused the diarrhea as the most common side effect ([6/30 (20.00%)]). Although panitumumab and its combination with bevacizumab and chemotherapy showed detrimental effects vs. 0.66%), vs. 1/602 (0.17%])

Comparison of the EGFR Mabs with VEGF Mabs and their combination for mCRC treatment

Bevacizumab acts as anti Vascular Endothelial Growth Factor (VEGF) and is used to inhibit VEGF function in vascular endothelial cells and inhibit tumor angiogenesis. Application of bevacizumab plus chemotherapy in most randomized controlled trials of mCRC patients showed a significant increase in the PFS or...
disease free survival rate 44. The combination of bev-
acizumab with oxaliplatin in another clinical trial result-
ed in better OS and PFS rate compared to the situa-
tion when panitumumab was added to bevacizumab and
oxaliplatin (PFS for Bmab+Oxaliplatin 11.4 months vs.
Bmab+Oxaliplatin+ Pnab 10 months, HR=1.27 CI;
95%, p=0.011) (OS for Bmab+Oxaliplatin 24.5 months vs.
Bmab+Oxaliplatin+ Pnab 19.4 months, HR=1.43,
CI, 95%, p=0.005) 44. The results of this study indicate
the superiority of bevacizumab plus chemotherapy than
both panitumumab and bevacizumab plus chemothera-
py. Bevacizumab beyond first progression is also associ-
ated with a longer median OS time and has an im-
portant role in improving the overall success of therapy
for mCRC patients 45. Application of FOLFOXIRI plus
bevacizumab in comparison to FOLFIRI plus beva-
cizumab resulted in better PFS rate as 12.1 months in
the first group and 9.7 months in the second group
(HR=0.75; 95% CI=0.62 to 0.90; p=0.003). OS was
also longer in the first group, but it was not significant
46. These results may imply to the benefits of the com-
bination of bevacizumab with oxaliplatin.

In a study to compare the panitumumab plus modi-
fied FOLFOX6 and bevacizumab plus modified
FOLFOX6 for the treatment of mCRC patients, the obtained
results showed similar PFS between the two
arms and median OS of 34.2 months and 24.3 months
in the panitumumab and bevacizumab arms, respec-
tively (HR, 0.62; 95% CI, 0.44 to 0.89; p=0.009). Re-
garding the results of pervious mentioned studies, al-
though the addition of panitumumab to the bevaci-
zumab and chemotherapy caused the OS to decrease,
thereby increasing the incidence of side effects 34,35, but
in comparison to bevacizumab, OS was significan-
tly improved when panitumumab was combined with
mFOLFOX6 in mCRC patients 47.

Santos et al have shown that in mCRC patients
treated with cetuximab or bevacizumab, the median
PFS was around 11.7 months in the cetuximab arm and
9.6 months for bevacizumab arm 48,49. In addition,
combination of FOLFIRI and cetuximab resulted in
prolonged overall survival in comparison to FOLFIRI
plus bevacizumab in first line treatment for mCRC
patients (22.7 vs. 18.7 months), although this differ-
ence was not statistically significant (HR=0.86, 95%
CI, 0.55-1.35; p=0.55) 50. Cetuximab also showed a
superiority in the first line treatment of mCRC patients
with KRAS p.G13D-mutation in comparison to beva-
cizumab as OS was 20.1 months in patients treated
with cetuximab and 14.9 months in patients receiving beva-
cizumab-containing regimens (hazard ratio: 0.70, p=
0.29) 51. In a phase II study of capecitabine, oxaliplatin,
and cetuximab with (arm A) or without (arm B)
bevacizumab, results showed an overall response rate
of 54% (36.4% in arm A and 72.7% in arm B). PFS in
this study was 8.7 months in arm A and 14.4 months
in arm B. Also, the median overall survival was 18.0
months in arm A and 42.5 months in arm B 52. Also in

Phase III trial, in order to study the effect of cetuximab
plus bevacizumab, and 5-fluorouracil/leucovorin in
comparison to FOLFOX-bevacizumab in mCRC pa-

tients, results showed that cetuximab plus beva-
cizumab, and 5-fluorouracil/leucovorin was not superior
to FOLFOX6-bevacizumab in terms of 12-month PFS
(45%/32%, respectively) and overall response rate
(52%/41%, respectively) 53. Regarding all mentioned
trials, cetuximab has more privileges to bevacizumab
since it can be combined with chemotherapy for mCRC
treatment, but is not an appropriate candidate in com-
bination with bevacizumab and in chemotherapy regi-
men and this combination is not beneficial for mCRC
treatment.

Aflibercept is another anti VEGF inhibitor that
binds to circulating VEGFs and acts like a VEGF trap,
inhibits the neovascularization in the choriocapillaris or
the tumor. Aflibercept in combination with 5-fluo-
ruoracil, leucovorin and irinotecan (FOLFIRI) signifi-
cantly improved survival in a phase III study of pa-

tients with mCRC who had a history of treatment with
oxaliplatin 54, albeit the outcome of aflibercept treat-
ment in mCRC patients mainly depended on the history
of previous treatment with bevacizumab. Patients who
received bevacizumab before aflibercept usually show-
ed a weak response 55. Due to lack of large clinical
trials, it is difficult to compare aflibercept efficacy and
safety as monotherapy and in combination with Mabs
in this field. But regarding the results of studies on
bevacizumab and its combination with cetuximab and
panitumumab, it can be concluded that combining
VEGF and EGFR inhibitors in metastatic colorectal
cancer is not always advantageous. However, any of
these agents can lead the treatment to success in combi-
nation with chemotherapy.

**Conclusion**

In many cases, cancer becomes metastatic before
correct diagnosis and makes the treatment difficult.
Although by introducing different chemotherapeutic
agents such as irinotecan and oxaliplatin in combina-
tion with fluorouracil or leucovorin, the hope for sur-
vival is created in patients with mCRC, most of these
patients fail to be cured due to the development of drug
resistance. This drug resistance urges researchers to
explore an alternative approach for chemo refractory
patients and for those who do not respond to conven-
tional chemotherapy.

Anti-EGFR antibodies can be considered as appro-
priate candidates for mCRC patients who have no re-

dponse to chemotherapy. Results of many clinical trials
showed the effectiveness of these Mabs in the treat-
ment of mCRC patients in different line settings and at
any progressive stage. However, it should not be ig-
nored that neither conventional chemotherapy nor anti-
EGFR therapy alone will not help mCRC patients, ra-
ther the best achievement is only accessible when anti-
EGFR Mabs are used in combination with routine
chemotherapy. For example, combination of mFOLF-OX6 and FOLFIRI with cetuximab resulted in a better objective response rate and overall survival in comparison to each chemotherapy regimen alone. Or in a PRIME phase III trial, administration of panitumumab plus FOLOX4 significantly increased the PFS in mCRC patients compared to FOLFOX4 alone.

An important factor in deciding to include the cetuximab or panitumumab in the treatment of mCRC patients is the KRAS mutation status. In fact, the selection of reliable patients before starting the interventions provides an opportunity to increase the response rate and chance of treatment success. This is a unique trait of anti-EGFR therapy. Also, despite high drug resistance to conventional chemotherapy which is considered as a main challenge, resistance to cetuximab and panitumumab among mCRC patients seems unlikely. The only superiority of routine chemotherapy agents in comparison to anti-EGFR inhibitors is their easy access due to their generic availability and lower price. In addition to anti-EGFR inhibitors, anti VEGF Mabs such as bevacizumab also showed effectiveness in the treatment of mCRC. But like cetuximab and panitumumab, bevacizumab also showed maximum effectiveness when combined to routine chemotherapy. Although some clinical trials compared the effects of bevacizumab and cetuximab and/or panitumumab, additional clinical trials are still required to evaluate the superiority of anti-EGFR or anti VEGF inhibitors. However, regarding what was reviewed in this article, anti-EGFR therapy does not have the ability to be replaced by chemotherapy.

The recommended dose and schedule for cetuximab is 400 mg/m² administered intravenously as a 120 min infusion as an initial dose, followed by 250 mg/m² infused over 30 min weekly. For panitumumab, recommended dose is usually 6 mg/kg every 14 days as an intravenous infusion over 60 min. The best treatment option is usually selected by considering factors related to both patients and clinics. Maximum effectiveness and minimum adverse effects are among these factors. Comparing cetuximab and panitumumab in many clinical trials, better overall survival is often achieved by cetuximab plus chemotherapy and better progression-free survival with panitumumab plus chemotherapy. Also, cetuximab in comparison to best supportive care can significantly improve the overall survival in the first line setting of mCRC treatment. However, application of cetuximab is not always associated with good results. In EPOC trial, cetuximab when combined with fluoropyrimidine and oxaliplatin showed the controversial results and even worsened the PFS. Panitumumab also helped chemotherapy to be more efficient. In many randomized clinical trials, significant improvements in tumor response rates and progression-free survival were observed when panitumumab was combined with chemotherapy or used as monotherapy in chemorefractory mCRC individuals. But in contrast to cetuximab, panitumumab mostly conferred PFS benefit. The positive fact about panitumumab is its ability to improve treatment outcome among previously cetuximab treated mCRC patients. Meanwhile, regarding the key role of KRAS status, it is fundamentally important to have an appropriate method for selection of patients through the use of novel predictive biomarkers to enhance the chance of treatment success.

The other important factor in treatment decision making is minimum adverse effects of therapy. Between cetuximab and panitumumab, there is no meaningful difference based on treatment related side effects. As mentioned before, most common adverse effects of anti-EGFR therapy are skin rash, electrolyte abnormality and infusion reaction. But some detrimental side effects such as lung infection and embolism and treatment related death were observed by the application of cetuximab. However, these observed side effects is not a justifiable evidence to consider cetuximab inferior in comparison to panitumumab. In contrast to the application of one biological drug from anti-EGFR and anti VEGF inhibitors, it was also observed that in combination therapy of bevacizumab based chemotherapy with cetuximab and/or panitumumab, the addition of panitumumab was associated with detrimental effects and it raised the incidence of side effects, while the incidence of side effects has been lowered when cetuximab was combined with bevacizumab based chemotherapy. Nevertheless, to establish the superiority of cetuximab or panitumumab for treatment of mCRC, clinical trials with larger sample size are needed. On the other hand, although fewer side effects are an advantage of any therapeutic modality, a noticeable fact about the application of anti-EGFR therapy is that more effectiveness is reported in those patients with more side effects. Regarding the results of mentioned clinical trials, the decision of treatment option between these two available anti-EGFR Mabs as stated before depends mostly on the patient condition and clinical access.

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