A glyco-engineered anti-HER2 monoclonal antibody (TrasGEX) induces a long-lasting remission in a patient with HER2 overexpressing metastatic colorectal cancer after failure of all available treatment options

Colorectal cancer (CRC) is one of the most common forms of malignancies and the second leading cause of cancer-related death in Western countries. Although chemotherapy in combination with novel antiangiogenic or antiepidermal growth factor receptor (EGFR)-directed agents has significantly improved the survival time of patients with metastatic CRC (mCRC), patients with metastatic disease still face a very poor 5-year survival rate less than 10%. The human EGFR 2 (HER2) has proven as an effective drug target in breast and gastric cancers. In CRC, data about the therapeutic utility of anti-HER2 agents are sparse, and had been only reported for treatment-naïve or second-line-treated patients. There is only one clinical phase II study reporting the use of the first-generation HER2/neu-directed trastuzumab in combination with irinotecan in first-line or second-line setting. Although some activity of this drug combination has been observed, the study was terminated early due to insufficient patient recruitment. In this patient case, we report about the successful use of a novel glyco-engineered anti-HER2 monoclonal antibody (TrasGEX) with 10-fold to

**Figure 1** Representative picture of HER2/neu immunohistochemistry of colorectal cancer tissue. (A) A strong brown (+3) reaction in the tumour tissue indicates HER2/neu overexpression. In contrast, the surrounding normal colon mucosa (*) stains negatively (×100). (B) Higher magnification (×200) showing the homogenous, membranous staining pattern of the tumour cell.
A 61-year-old female Caucasian patient was newly diagnosed with CRC in December 2002, and consequently underwent a left hemicolectomy at the Department of Surgery, Medical University of Graz. A stage II pathological T4N0 CRC was found according to the tumour, node, metastases classification system. One month later, a single liver metastasis was diagnosed by CT, and pre-operative chemotherapy comprising eight cycles of FOLFIRI/bevacizumab was initiated. Five months later, a partial remission of the liver metastasis was achieved, and a resection of the liver metastasis (segment VI) was performed in July 2003. After a 30-month disease-free period and routine clinical follow-up, a CT scan of the abdomen detected enlarged retroperitoneal lymph nodes. Based on an individual patient decision and treatment approach, the patient underwent lymph node resection and post-operative chemotherapy with seven cycles of FOLFOX4 regimen. Oxaliplatin had to be dose-reduced after two cycles due to a prolonged grade III leucopenia. Another 9 months later, the disease recurred with a solitary liver metastasis in segment VI and one single metastasis in the right suprarenal gland. The patient again underwent liver resection and subtotal right adrenalectomy in April 2007. Three months after surgery, solitary retroperitoneal lymph node metastases were detected and consecutively resected. After a 4-month disease-free period, a recurrence of metastasis in the liver was radiologically detected. Palliative chemotherapy with FOLFIRI/bevacizumab was administered for eight cycles, resulting in partial remission of the liver metastasis.

Unfortunately, half a year after a break of the therapy, the patient experienced a disease progression with spread to multiple metastases in both lungs and solitary liver metastasis. Rechallenge of FOLFIRI/bevacizumab (80% dose reduction of irinotecan due to prolonged haematotoxicity) resulted again in partial remission after 12 cycles. In February 2011, the disease progressed again, and after determining KRAS exon 2 mutational status (a KRAS wild-type was diagnosed by pyrosequencing in the tumour tissue, a palliative therapy containing irinotecan (150 mg/m²) plus panitumumab (6 mg/kg) was administered every 2 weeks for five cycles. Despite the use of this combination therapy and the administration of another two palliative therapy approaches with FOLFOX/bevacizumab (4 cycles) as well as UFT (tegafur 2-1-1 day 1–28, leucovorin 2-2-2 day 1–28 for 4 months), no objective response or clinical benefit could be reached. A further molecular characterisation of the tumour tissue detected no mutations in KRAS/NRAS exons 3 and 4, as well as no BRAF V600E mutation. Facing the disease progression and the use of all currently available drugs, the patient consented to participate in the phase I TrasGEX clinical trial (ClinicalTrials.gov number NCT01409343). After consenting, a HER2/neu immunohistochemical expression in the tumour tissue was evaluated in March 2012 (Dako HercepTest). Around 95% of tumour cells showed a strong complete membranous staining pattern (3+), leading to the enrolment of the patient into the TrasGEX dose escalation protocol (figure 1A, B).

TrasGEX is a humanised IgG1 anti-HER2 antibody, and belongs to a group of antibodies, which are glyco-engineered by the GlycoExpress platform with a fully human glycosylation and optimised key sugar determinants, including fucose, galactose, sialylation and branching. Major optimisation for TrasGEX is an enhanced ADCC antitumour activity in combination with other variable domain-mediated antitumour activities. ADCC is mediated via the FcRIIIa receptor on immune effector cells. Two allotypes of this receptor differing in amino acid position 158 (V158F) are known, which have different affinities to human IgG1, depending on the glycosylation of the glycans in the IgG1 Fc tail. This difference in receptor affinity results in a reduced natural killer cell activation for F-containing allotypes for antibodies generated in Chinese hamster ovary (CHO) or other rodent production systems and, therefore, in a reduced ADCC activity.

Figure 2  (A and B) Baseline imaging before therapy with TrasGEX has been initiated. (A) Multiple lung metastatic lesions with one target lesion of 2 cm in diameter (red lines) and (B) a liver metastasis with 11 cm in the largest diameter. (C and D) Three cycles later, the patient has a decrease of size in lung (C) and liver metastases (D). (E and F) Imaging 21 months (30 cycles) after initiation of TrasGEX. The target lung lesion diminished (E) and the target liver lesion measures 4.5 cm (F).
Patients with FcγRIIIa allotype (V/V) treated with trastuzumab (Herceptin) or cetuximab (Erbitux) have a better clinical outcome than those with F/F or F/V.6–8

According to the study protocol, the patient received 480 mg of TrasGEX intravenously over 4 h and every consecutive cycle on day 21 with an infusion time of 90 min. Three cycles later, the patient had a decrease of size in lung and liver metastases (sum of the longest diameters, target lesions in the liver and right lower lung lobe reduced from 19.7 to 11.1 cm (figure 2A, C), target lesion in the right liver lobe from 10 to 5.7 cm) (figure 2B, D), so, continuation of the targeted therapy according to the study protocol was performed. Until now, 21 months (30 cycles) after initiation of TrasGEX, the patient still has a partial remission of the disease (target liver lesion 4.5 cm), and presents in excellent clinical condition (figure 2E, F). Remarkably, over the whole study period, no side effects including cardiotoxicity (the left ventricular ejection fraction determined by echocardiography remained unchanged throughout the 21 months) occurred.

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