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## N&WICS - TA439 - Cetuximab - Previously untreated metastatic colorectal cancer

Before providing patient identifiable data on this form, please confirm that the patient (or in the case of a minor or vulnerable adult with the parent/legal guardian/carer) has given appropriate explicit consent for sensitive personal information on this form to be passed to the CCG and/or

| CSU for processing this funding red  | uest and validating subsequent in  | nvoices. Consent given:      |                        |                    |           |
|--|--|------------------------------|------------------------|--------------------|-----------|
| If there is more than one NICE-app<br>about the advantages and disadvar<br>patient is likely to adhere to treatm   | tages of the treatments available  | . This has taken into cons   | ideration therapeu     | tic need and wheth | ner or no |
| and price per dose) unless an orde   | of preference is stated in the TA  | s. 🗌                         |                        |                    |           |
| Patient<br>NHS No:   | Trust:   |                              | Practice<br>Name:      |                    |           |
| Patient<br>Hospital<br>No:   | Consultant Making Request:   |                              | Practice<br>Postcode:  |                    |           |
| Patient's<br>Initials and<br>DoB:  |  |                              | Practice<br>Code:      |                    |           |
| Notification Email Address:  | (@NHS.net a  | ccount ONLY)                 | Contact name & number: |                    |           |
| Start date of requested treatment:   | Provider: Private [ Supplier: Homecal                                    | □NHS<br>re □Hospital         | Sub-Type:              | /A 🔽               |           |
| By completing this form, you confir<br>commissioning statement. Any requ<br>For support regarding IFRs, please<br>For support regarding the criteria lie   | uests which fall outside of this use contact: norfolkicd@nhs.net         | e will require an individual |                        |                    | al        |
| Please indicate whether patie  | nt meets the following NICE c  | riteria:                     |                        | Please tick        |           |
| epidermal growth factor receptor in combination with:  | within its marketing authorisation, (EGFR)-expressing, RAS wild-ty       |                              |                        | □Yes □No           |           |
| 5-fluorouracil, folinic acid and oxaliplatin (FOLFOX) or     5-fluorouracil, folinic acid and irinotecan (FOLFIRI).  |  |                              |                        |                    |           |
| Panitumumab is recommende  | d, within its marketing authorisatictal cancer in adults in combinati    |                              | ously untreated        |                    |           |
| • FOLFOX or  |  |                              | ☐Yes ☐No               |                    |           |
| • FOLFIRI.   |  |                              |                        |                    |           |
| 3. The drugs are recommended only when the companies provide them with the discount agreed in the patient access scheme (for panitumumab) or commercial access agreement (for cetuximab).  |  |                              | ☐Yes ☐No               |                    |           |
|  | ent should normally be started wi<br>e and any additional administration |                              | ug (considering        |                    |           |
| Please confirm that the choice of drug is considered the most cost-effective treatment for this individual patient, noting that alternative treatment options recommended by NICE may be lower in cost, including the use of biosimilar medications if applicable. |  |                              | ☐Yes ☐No               |                    |           |
|  | treatment is available and has no cribing has been included in the       |                              |                        |                    |           |

5. The product is being used as described by local commissioning position.

☐ Yes ☐ No

| I confirm that the patient meets the criteria for treatment |                  |
|---|------------------|
| Name of person completing:                                  | Contact Details: |
| Designation of person completing:                           | Date:            |
| Trust Authorising Pharmacist                                |                  |
| Name:  Date:  |                  |