What carcinoembryonic antigen level should trigger further investigation during colorectal cancer follow-up? A systematic review and secondary analysis of a randomised controlled trial

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Scientific summary

Follow-up of the FACS RCT

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Scientific summary

Background
Following primary surgical and adjuvant treatment for colorectal cancer, patients are routinely followed up with blood carcinoembryonic antigen (CEA) testing for 5 years. The Follow-up After Colorectal Surgery (FACS) trial showed that this follow-up is effective at detecting recurrences treatable with curative intent. However, the optimal testing interval and method for interpreting test results lack a firm evidence base. Our initial protocol was restricted to conducting a secondary analysis of CEA testing results from the FACS trial. However, initial work revealed serious limitations of existing reviews of previous research and so we also conducted a formal systematic review following Cochrane guidelines for identifying and meta-analysing studies of diagnostic accuracy.

Aim and objectives
The main aim was to determine how the CEA test result should be interpreted to inform the decision to undertake further investigation to detect treatable recurrences. Secondary objectives were to determine whether or not diagnostic accuracy could be improved by (1) taking account of the baseline CEA level and other pretest patient characteristics; (2) considering the trend in CEA levels over time; (3) considering whether recurrence occurred early or late in follow-up; and (4) changing the testing interval.

Methods for the systematic review

Search
The search details are reported on the Cochrane website [http://onlinelibrary.wiley.com/doi/10.1002/14651858.CD011134.pub2/full (last accessed 6 November 2016)].

Studies included
Cross-sectional diagnostic test accuracy studies, cohort studies or randomised trials, conducted in primary care or hospital settings, involving adults with no detectable residual disease after curative surgery (with or without adjuvant therapy) and reporting results extractable in a 2 × 2 format (i.e. test +/− by case +/−).

Index test
The index test was the blood CEA level.

Reference standard
The reference standard was clinical diagnosis of recurrence of colorectal cancer following primary treatment confirmed by imaging, histology or clinical follow-up [for details of the reference standard see the full report on the Cochrane website: http://onlinelibrary.wiley.com/doi/10.1002/14651858.CD011134.pub2/pdf (accessed 25 August 2016)].

Analysis
Two review authors extracted data independently and three authors independently performed a Quality Assessment of Diagnostic Accuracy Studies 2 assessment of the included studies, with subsequent discussion to reach consensus on overall judgements of risk of bias and applicability. The meta-analyses followed Cochrane guidelines for pooling test accuracy data.
Methods for the main analysis

Design
This study involved a secondary observational analysis of data from the FACS trial, a 2 × 2 pragmatic randomised factorial controlled trial comparing minimum post-surgery follow-up of colorectal cancer patients for 5 years with 3- to 6-monthly blood tests for CEA and 6- to 12-monthly computerised tomography (CT) imaging. We analysed the two arms of the trial that required CEA testing.

Statistical analysis
Receiver operating characteristic (ROC) analysis was implemented to compare the effect of making the decision to trigger further investigation in individual patient on the basis of (1) the CEA level at each test; (2) the difference between test and postoperative CEA levels (expressed both as an absolute value and as a ratio); and (3) the trend in CEA levels over time. An operational analysis of the probable impact of CEA testing if used prospectively in clinical practice was also conducted, hypothetically applying the four most commonly reported thresholds in the systematic review (2.5, 5, 7.5 and 10 µg/l) to trigger further investigation on the basis of each test carried out during the follow-up period. To investigate the diagnostic accuracy of assessing trends in serial CEA measurements within an individual rather than simply interpreting the most recent CEA measurement taken, linear regression models were fitted to the CEA values for each individual over time. The distribution of slope coefficients for individuals who did and did not experience recurrence were compared and ROC analysis was used to evaluate the diagnostic accuracy of CEA trend. All analyses were carried out using the statistical package R [see www.R-project.org/ (accessed 25 August 2016)]; the ROC analysis was carried out using the R package pROC.

Participants
Patients who had undergone curative surgery for primary colorectal cancer and who, after extensive testing (histology, imaging and a CEA level of ≤ 10 µg/l), were confirmed to have no residual disease were recruited from 39 NHS hospitals across all regions of England. The analysis was based on 582 patients from the two arms of the study that received CEA testing.

Carcinoembryonic antigen measurement
The CEA analysis was undertaken using a Siemens Centaur XP analyser (Siemens Healthcare, Erlangen, Germany) at a single laboratory with a standard quality control regime to ensure longitudinal stability. If the blood CEA level was ≥ 7 µg/l above the patient’s baseline level at trial entry after repeat measurement, the general practitioner was asked to refer the patient urgently to the local hospital for further investigation. The median number of CEA measurements available for each participant was 13 [interquartile range (IQR) 10–14], with a median of six (IQR 3–9) measurements in patients who developed a recurrence and 14 (IQR 13–14) in those who did not develop a recurrence.

Diagnostic reference standard
The reference standard against which diagnostic accuracy was assessed was clinical diagnosis of recurrence of colorectal cancer as determined by the colorectal cancer multidisciplinary team at the participating hospital centre.

Results
Diagnostic accuracy of a single test
The diagnostic accuracy of CEA testing across all thresholds, estimated on the basis of all CEA tests carried out prior to diagnosis, is modest [area under the receiver operating characteristic curve (AUC) 0.74, 95% confidence interval (CI) 0.68 to 0.80]. Sensitivity is estimated as 50.0% (95% CI 40.1% to 59.9%). The median lead time gained at a recommended threshold of 5 µg/l is about 3 months, but the predictive value would be 62% assuming the same frequency of recurrence experienced in the trial, implying that about four in 10 patients without a recurrence will have at least one false alarm. The positive predictive value of
an individual test (rather than an individual patient) is even lower at 43.3% (95% CI 35.8% to 51.0%), as some patients suffer repeated false alarms. For example, the 89 false alarms triggered at a threshold of 5 µg/l were clustered in 29 individuals, 15 of whom (51.7%) would have more than one false alarm and eight of whom would have more than five false alarms. Trying to improve the sensitivity of CEA testing by reducing the threshold for further investigation has a high cost in terms of falling specificity. Although sensitivity can be increased to 63.5% (95% CI 54.2% to 72.8%) by reducing the threshold to 2.5 µg/l, there is a sevenfold increase in the number of times further investigation is triggered and, in 84% of cases, no recurrence is detected.

**Adjusting for postoperative baseline carcinoembryonic antigen level**

Adjusting the CEA level by an individual’s baseline measurement offers no notable improvement in diagnostic accuracy. Of the 6623 CEA measurements in the database, 3881 (59%) were lower than their baseline measurement and therefore had a negative ‘difference’ value or a ratio value of < 1; 19 patients who developed recurrence always had a negative adjusted value (i.e. all of their CEA measurements were lower than their baseline level).

**Predicting missed cases and false alarms**

None of the characteristics assessed (patient age and smoking status, site and stage of the primary tumour, receipt of adjuvant therapy and delay in commencing monitoring and site of recurrence) significantly increased the likelihood of recurrence being missed. However, current smoking was significantly predictive of multiple false alarms (adjusted odds ratio 6.55, 95% CI 1.52 to 28.20; \( p = 0.01 \)).

**Diagnostic accuracy of trend**

The AUC suggests that the rate of change of CEA level provides better overall discriminatory power than the single-value CEA transformations explored (AUC for positive trend 0.85, 95% CI 0.78 to 0.91). A negative trend (i.e. a reducing level of CEA post treatment) may also have diagnostic value, increasing the AUC to 0.91; however, this improvement is not statistically significant. The optimal threshold for interpreting trend changes over time, to achieve 70% sensitivity at around 90% specificity, falls from 1.75 µg/l in year 1 to 0.3 µg/l in year 5.

**Diagnostic accuracy in early and late recurrence**

Assessing trend performs better than single test assessment in detecting both early recurrence (recurrence in the first 2 years after treatment) and late recurrence (recurrence in years 3–5), with little difference in accuracy for each time period.

**Test interval**

At a single test threshold of 5 µg/l, the testing interval needs to be approximately halved in year 1 to ensure that the number of recurrences detectable at each test, and, therefore, test operational performance, remains fairly constant over time. The test interval also needs to be reduced in year 1 if action is to be taken on the basis of trend [with a 3-monthly testing interval the CEA level would not be measured with any precision until month 9, by which time 31 (29.8%) recurrences had already been diagnosed].

**Relevance and implications**

**The importance of not triaging with carcinoembryonic antigen alone**

Our main analysis confirms the findings of the systematic review: CEA testing alone is insufficient as a triage test for colorectal cancer recurrence. Whatever threshold is applied to interpret the CEA test result (based on a single test or trend), a significant number of patients will suffer recurrence without a detectable change in CEA levels. This underlines the importance of combining CEA testing with scheduled imaging, as recommended in most national guidelines.
The advantage of making decisions on the trend in carcinoembryonic antigen levels

The diagnostic accuracy of the trend in CEA levels, assessed by the slope (beta-coefficient) of the linear regression line, was consistently better than interpreting the results of a single test, regardless of whether the single test was adjusted for the baseline postoperative CEA level. The observation that optimal performance was achieved by taking account of a negative as well as a positive trend merits further investigation. It suggests that a slow post-treatment reduction in CEA level is itself a marker of recurrence.

The choice of carcinoembryonic antigen threshold

Both the systematic review and the main analysis highlighted the very high cost in terms of false alarms of adopting an action threshold for a single test below the 5 µg/l commonly recommended by national guidelines. The analysis of operational performance suggests that a higher threshold (of perhaps 10 µg/l) may be preferable. Even using trend analysis, the number of false alarms suggests that aiming for a sensitivity of 70% – augmenting CEA testing with a colonoscopy and one or two CT scans to detect the missed 30% of recurrences – may be the clinically preferable option. In applying the trend analysis, the main results also highlighted the importance of not applying the same action threshold throughout the 5-year follow-up period. It is important that thresholds derived from our data are checked and refined in an experimental setting before being rolled out, but, as a starting point, we suggest initiating further investigation if the rate of change in CEA level exceeds 1.7 µg/l/year in year 1, 1.4 µg/l/year in year 2, 0.8 µg/l/year in year 3, 0.5 µg/l/year in year 4 and 0.3 µg/l/year in year 5.

The choice of testing interval

The testing frequency in year 1 would need to be increased to achieve a more consistent level of operational performance over time and to allow for an earlier assessment of trend. A testing schedule of monthly for the first 3 months and then every 2 months for the rest of the year would be consistent with our findings. Adopting this increased testing frequency would be challenging in some health-care systems and would need careful planning (as it requires rapid turnaround of results, good communication with patients and access to a clinician who is able to discuss and act on the results quickly). Although the falling incidence of recurrence would suggest that testing frequency should be reduced to one test in year 5, this would have implications for the achievable lead time.

Who should and should not be followed up with carcinoembryonic antigen

The main analysis of the FACS trial shows that CEA follow-up is appropriate for all patients who have completed surgical and adjuvant treatment for their colorectal cancer and who have, on extensive investigation, no sign of recurrence. It also shows that patients at Dukes’ stages A–C have a similar incidence of treatable recurrence and obtain similar benefit. There is no suggestion that patient age, the characteristics of the primary tumour or the recurrence site predict either missed cases from non-response or false alarms. However, the likelihood of multiple false alarms is significantly higher in smokers, suggesting that CEA is not an appropriate follow-up method for patients who continue to smoke.

Other research implications

The systematic review drew attention to the poor quality of the majority of diagnostic studies on CEA testing. Moreover, virtually all studies assessed CEA as a single diagnostic test, ignoring the fact that it is used as a monitoring test and is carried out repeatedly over time. Even the Cochrane diagnostic accuracy methodology that we used to conduct the systematic review considers only single test results, not trend over time. This issue should be addressed, not just in relation to CEA testing but in relation to all tests that are repeated over time to monitor disease progression. The need to refine the suggested cut-off points for monitoring CEA trend (by conducting pilot implementation studies before large-scale roll-out) has already been mentioned. In conducting these studies, it might be preferable to use a reference standard based on recurrence treatable with curative intent rather than any recurrence.
Study registration

This study is registered as PROSPERO CRD42015019327 and ISRCTN93652154.

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