Rezumat

Recurențele locale (RL) ale cancerului de rect (CR) au o rată de supraviețuire în afara bolii de până la 50% dacă sunt diagnosticate devreme. Ultrasonografia endorectală (USER) combinată cu utilizarea tușelui rectal sunt două metode de urmărire a pacienților care pot fi realizate într-un cabinet de consultație. Acesta este primul studiu realizat pentru a determina accuratețea diagnostică a USER în detectarea RL din CR și a determina dacă reprezintă o metodă de încredere în supravegherea pacienților. Trei autori au căutat independent în bazele de date MEDLINE și ClinicalTrials.gov și au inclus articolele originale relevante bazat pe criteriile includere/excludere. 3220 de articole au fost identificate inițial, din care 50 au fost selectate după citirea abstractelor. 22 de articole au fost incluse în final, reprezentând 3737 de pacienți supravegheați 59,72 ± 16,4 luni. Bazat pe datele disponibile, sensibilitatea USER a fost 88,3% (CI 84,6 – 91,3%), specificitatea egală cu 94,3% (CI 92,7 – 95,5%) și rata relativă de diagnostic a fost 271,88 (CI 76,998 – 960,04). USER a fost singura metodă care a diagnosticat RL în 40 ± 12 % (CI 28 – 52 %) din cazuri. Suprafața de sub curba sROC a fost 0,9723 ± 0,131. După tratamentul curativ al CR au apărut RL la 15 ± 2,99% din pacienți. În concluzie, USER pare să fie o metodă bună și eficientă de supraveghere a pacienților pentru depistarea RL după CR.
Introduction

Locally recurrent rectal cancer is characterised by clinical, radiological and/or pathological determination of rectal cancer recurrence in the pelvic region, following a curative treatment (1). Even though, in the last three decades, the incidence of local recurrence (LR) has dropped from 30% to less than 10%, LR still affects a large number of oncologic patients which suffer from symptoms such as: localized pelvic pain, bleeding, fistula and, ultimately, death (2). With the development of chemotherapy, radiotherapy and “watch-and-wait” strategy, the early detection of LR is of utmost importance in the follow-up of rectal cancer patients, owing to better survival and better functional outcomes (3). Early LR can be treated either by salvage surgery or by local excision depending on the stage of the LR at diagnosis, leading to a disease-free survival rates ranging between 20 to 50 percent (4-8). Thus, a good follow-up strategy has an important role in the management of the patient post-treatment.

Endorectal ultrasound or transrectal ultrasound (ERUS/ TRUS) is an outpatient procedure that uses an ultrasound probe introduced in the rectal lumen in order to visualise the rectal wall and its layers. Compared with MRI, CT or digital examination only, ERUS has been shown to be a better choice in pre-operative staging of local rectal cancer to determine local tumour invasion (9). In the same meta-analysis (9), neither of the procedures had a higher accuracy at staging lymph node involvement. Even though, there is still no agreement on what imaging method has the highest accuracy in detecting early LR, digital examination combined with ERUS makes it an easy, safe and cheap method to follow-up with patients.

The aim of this systematic review is to determine the diagnostic test accuracy of...
ERUS/ TRUS in the detection of LR after rectal cancer and if it should be trusted as good follow-up method.

Material and Method

Literature search

A comprehensive systematic literature search was performed in the MEDLINE PubMed database and ClinicalTrials.Gov to identify published articles in English or enrolled clinical trials based on human subjects. The work done for this article was in accordance to the PRISMA 2020 Checklist. Relevant articles for follow-up after rectal cancer and the use of ERUS/ TRUS in the diagnosis of LR after rectal cancer were included. The MEDLINE search spanned between January 1986 and January 2021 and included the MeSH (Medical SubHeadings) terms: “transrectal ultrasound”, “transrectal ultrasonography”, “transrectal echography”, “endorectal ultrasound”, “endorectal ultrasonography”, “endorectal echography” AND “rectal cancer”, “rectal neoplasms”, “colorectal cancer”, “colorectal neoplasm”. To identify additional relevant reference, the reference list of included articles was checked manually for other studies of importance. For ClinicalTrials.Gov there were two independent broad searches based on a simple MeSH term: “recurrent rectal cancer” and “endorectal ultrasound”.

Data Extraction

The studies were reviewed by two authors working independently (AA; LIS;) There was a two-parts process consisting of reviewing the titles and abstracts of all articles in order to include them to a full-text review and reading the final full-text analysis of the selected articles, eliminating from the review the ones that do not follow our criteria. In order to assure a 90% inter-reviewer similarity, the two authors swapped and compared the databases to check variables. Whenever it was necessary, a third reviewer (FDU) was called upon to decide including or excluding studies if the first two did not agree. The authors extracted the following data whenever it was available: author, year, funding, study design, location, number of centres, type of study (therapeutic or diagnostic), true positives, false positives, false negatives, true negatives, number of patients, percentage of patients diagnosed by ERUS/ TRUS alone, local recurrence rate, time of follow-up (FU) (maximum time period mentioned for each study), NCT registration and conclusion.

Study Selection

All included studies are in-line with the following inclusion criteria: (a) either a retrospective, prospective or randomized controlled trial; (b) more than 25 patients; (c) including follow-up of rectal cancer or colorectal cancer patients or diagnosis of LR using ERUS/ TRUS; (d) full-text available in English language. All articles that were: (a) reviews, editorial letters, animal studies, treatment protocols, diagnosis protocols, epidemiology studies or (b) studies that did not include ERUS/ TRUS were excluded. Some articles that followed all of the above criteria were excluded because they did not provide sufficient data about the follow-up of patients and no data could be extracted regarding diagnosis accuracy or the use of ERUS/ TRUS.

Statistical Analysis

Whenever possible, the data was extracted directly from the published articles. The measured outcomes were sensitivity, specificity, diagnostic odds ratio (DOR), area under the curve (AUC), percentage of LR diagnosed only by ERUS/ TRUS and local recurrence rate. Authors define sensitivity as the probability that ERUS/ TRUS will be positive in local recurrence cases and specificity as the probability that ERUS/ TRUS result will be negative in a case without local recurrence (10). DOR is defined as a single number that describes how many times higher are the chances of obtaining an ERUS/ TRUS positive result in LR than in a patient without LR
and it summarizes the diagnostic accuracy of ERUS. We also used Youden's Index as a general index of ERUS/TRUS accuracy and was calculated as sensitivity + specificity – 1 (10). Summary receiver operating characteristics (sROC) and AUC were developed and calculated. In general, an AUC close to 1 means ERUS/TRUS is the best test for LR after RC and a value close to 0.5 would mean ERUS/TRUS is a poor test for LR after RC. Q* is the intercept of the sROC and antidiagonal line through the unit square. Q* is used to indicate the overall accuracy of a test by comparing the case if sensitivity and specificity are the same, resulting that the closer the Q* to 1, the better the accuracy. In order to take into consideration, the inconsistency and heterogeneity of studies between the pooled time intervals we used I² statistics to calculate the percentage of variation across studies that is due to heterogeneity rather than chance (11, 12) with values closer to 0% meaning the confidence intervals of data across studies closely overlaps, whereas a value larger than 50% means there is low overlap between confidence intervals among studies. Usually, values of I² have 25%, 50% and 75% cut-offs for low, moderate and high inconsistency. Authors also looked at likelihood ration that ERUS/TRUS would be expected in a patient with LR compared to the likelihood of the same result to be seen in a patient without LR (10). Confidence intervals (CI) were set at 95%. Authors also tried to explore sources of heterogeneity in the studies using Moses-Shapiro-Littenberg meta-regression (multi-variate regression of the summary estimate DOR) (13) method of Weighted Least Squares by inverse variation (14). Data analysis was performed using RevMan 5.4.1 (The Nordic Cochrane Centre, Copenhagen, Denmark), MetaDisc 1.4 (Unit of Clinical Biostatistics Team of the Ramon y Cajal Hospital, Madrid, Spain) and IBM SPSS Statistics v27 (Chicago, Illinois, USA).

Results

After the search was performed there were 2806 articles identified from PubMed/Medline database and 414 clinical trials identified on ClinicalTrials.gov (363 trials for “recurrent rectal cancer” and 51 trials for “endorectal ultrasound”). After reading the abstracts and extensively cross-checked reference list of selected articles, authors selected 50 articles for full-text analysis and no clinical trial on NCT database was found to study the follow-up of rectal cancer with ERUS/TRUS or the diagnosis of LR after rectal cancer using ERUS/TRUS. Of the 50 articles selected, authors considered eligible for study inclusion only 22 articles. Reasons for not including studies were: (a) 11 articles were not available in English; (b) 6 articles considered recurrent rectal cancer an exclusion criteria; (c) 5 articles had no clear data to assess follow-up method; (d) for 3 articles authors did not succeed in finding a full-text article; (e) 1 article had fewer than 25 patients in follow-up population; (f) 1 article was studying endoscopic ultrasound; (g) 1 article did not differentiate among pelvic tumours (see Fig. 1).

418 data sets were retrieved from the 22 articles. 72.72% of the articles (16 articles) had a prospective study design, 27.28% had a retrospective study design (6 articles) and no article was a randomized controlled trial. Almost all articles were unicentric, with the exception of one article (15). Only 9.09% of studies (2 articles) (16,17) acknowledged their funding, both of them having no conflict of interest. The main focus of the included articles was follow-up (77.27%; 17 articles) and not treatment (21.74%; 5 articles). There were 3737 patients included in our database of 22 articles. Patients were followed for a maximum of 59.72 ± 16.4 (CI 43.3 – 76.1) months (see Table 1). In agreement with our search of literature in ClinicalTrials.gov, only one of the articles had a protocol published in a clinical trial registry (DRK) (16).

All included articles used ERUS/TRUS as a follow-up method as per our inclusion criteria. Further analysis of the literature for follow-up method included in at least one of the visits showed the following: digital examination (90.09%; 20 articles), biomarkers
(primarily CAE) (59.09%; 13 articles), MRI or CT (59.09%; 13 articles), colonography (50%; 11 articles) and PET-CT or scintigraphy (4.54%; 1 article) (see Table 5, Fig. 3). Only 16 articles had available data for meta-analysis. Sensitivity of ERUS/TRUS calculated from the pooled studies was 88.3% (CI 84.6 – 91.3%) with an inconsistency among studies of $I^2 = 68.3\%$ (see Table 4 and Fig. 2). Specificity of ERUS/TRUS was calculated for the selected studies at 94.3 % (CI 92.7 – 95.5%) with an inconsistency among studies of $I^2 = 90.0\%$ (Table 3 and Fig. 2). Positive likelihood ratio is 21.184 (CI 8.265 – 54.298) ($I^2 = 91.8\%$) meaning it was approximatively 21 times more likely to have LR if suspected findings arise during ERUS/ TRUS and negative likelihood ratio is 0.134 (CI 0.082 – 0.22) ($I^2 = 50.6\%$) meaning that only 13% of suspected lesions detected with ERUS/TRUS were of non-malign origin (see Table 5, Fig. 3). DOR of ERUS/ TRUS for the pooled studies was 271.88 (CI 76.998 – 960.04) ($I^2 = 69.6\%$) with an estimate variance for between-study of 4.0238 (see Table 6, Fig. 4). ERUS/ TRUS was the only diagnosis method to detect LR in 40 ± 12 % (CI 28 – 52 %) of detected LR (data available in 14 articles). The calculated rate of local recurrence after curative treatment of rectal cancer in our study was 15 ± 2.99 % (CI 12 – 18%) (data available from 22 articles) (see Table 3).

Developing sROC Curve (1-specificity), calculating AUC and $Q^*$ authors obtained a value for AUC = 0.9723 ± 0.0131 and $Q^* = 0.9235 ± 0.0219$ (see Fig. 3). This value and the aspect of sROC curve suggest there is no diagnostic threshold effect in our analysis (13). To further investigate this hypothesis, Spearman correlation coefficient between sensitivity and specificity was calculated using Moses’s model (18) and it is equal to -0.425 ($p=0.101$) which proves that there is no threshold effect.

In order to investigate for sources of heterogeneity (14), local recurrence rate and the period of follow-up were used in a meta-regression using extended Moses’s model (18) for covariates. The $p$-value of this analysis was 0.6122 for local recurrence rate and 0.5132 for follow-up period with CI including a 0 value for RDOR (relative DOR). This translates that the observed heterogeneity is not due to these specific covariates. Table 7 summarises the results of this diagnostic test accuracy meta-analysis on ERUS/ TRUS as a method of surveillance for LR after RC.

The authors have analysed whether the included studies had reached a positive, negative or neutral/undecided outcome. Of the 17 articles that looked at ERUS/ TRUS as follow-up method, 76.47% of studies (13 articles) reached a positive outcome, meaning that ERUS/ TRUS was considered a good follow-up method. 23.52 % of studies (4 articles) could not decide or did not have confidence in ERUS/ TRUS as follow-up.
method used by itself and no article reached a negative outcome, excluding ERUS/ TRUS from the follow-up protocol.

Discussion

In this systematic review authors summarized all relevant literature available to see whether ERUS/ TRUS is an efficient method of detection of LR after rectal cancer. ERUS/ TRUS was found to have a diagnostic odds ratio of almost 270 and an AUC = 0.973 which are strong indicators that ERUS/ TRUS is a fine tool in the diagnosis of LR after RC and was the only method able to detect LR in more

Table 1. Study design model, number of centres, type of study, number of patients and maximum length of follow-up (months) for the 22 included studies

| Article                        | Design (retrospective, prospective, randomized controlled trials); | No. Centres (unicentric, multicentric); | Group (Therapy study, Follow-up study) | No. Patients | Length of FU (months) |
|-------------------------------|---------------------------------------------------------------|-----------------------------------------|----------------------------------------|--------------|-----------------------|
| Kwakye et al, 2019 (38)       | retrospective                                                | unicentric                              | Follow-up study                        | 114          | 178                   |
| Manegold et al, 2019 (16)     | retrospective                                                | unicentric                              | Therapy study                          | 88           | 60                    |
| Li et al, 2015 (15)           | prospective                                                 | multicentric                            | Therapy study                          | 122          | 108                   |
| Park et al, 2015 (39)         | retrospective                                                | unicentric                              | Therapy study                          | 593          | 84                    |
| Marken et al, 2006 (21)       | retrospective                                                | unicentric                              | Follow-up study                        | 525          | 56                    |
| Hernandez de Anda et al, 2004 (22) | retrospective                                        | unicentric                              | Follow-up study                        | 275          | 61                    |
| Drudi et al, 2001 (24)        | prospective                                                 | unicentric                              | Follow-up study                        | 43           | 60                    |
| Hunerbein et al, 2001 (25)    | prospective                                                 | unicentric                              | Follow-up study                        | 312          | 60                    |
| Garcia-Aguilar et al, 2000 (4) | retrospective                                                | uncenter                                | Therapy study                          | 82           | 98                    |
| Lohnert et al, 2000 (40)      | prospective                                                 | unicentric                              | Follow-up study                        | 338          | 123                   |
| Ross et al, 1999 (41)         | retrospective                                                | unicentric                              | Therapy study                          | 272          | 50                    |
| Novell et al, 1997 (42)       | prospective                                                 | unicentric                              | Follow-up study                        | 140          | 44                    |
| Rotondo et al, 1997 (43)      | prospective                                                 | unicentric                              | Follow-up study                        | 62           | 26                    |
| Hunerbein et al, 1996 (44)    | prospective                                                 | unicentric                              | Follow-up study                        | 163          | 24                    |
| de Gara et al, 1995 (36)      | prospective                                                 | unicentric                              | Follow-up study                        | 30           | 24                    |
| Ramirez et al, 1994 (17)      | prospective                                                 | unicentric                              | Follow-up study                        | 66           | 60                    |
| Romano et al, 1993 (45)       | prospective                                                 | unicentric                              | Follow-up study                        | 37           | 25                    |
| Scalpi et al, 1993 (37)       | prospective                                                 | unicentric                              | Follow-up study                        | 35           | 24                    |
| Dressing et al, 1990 (46)     | prospective                                                 | unicentric                              | Follow-up study                        | 106          | 56                    |
| Beynon et al, 1989 (47)       | prospective                                                 | unicentric                              | Follow-up study                        | 83           | 54                    |
| Mascagni et al, 1989 (48)     | prospective                                                 | unicentric                              | Follow-up study                        | 120          | 15                    |
| Hildebrandt et al, 1986 (49)  | prospective                                                 | unicentric                              | Follow-up study                        | 129          | 24                    |

Table 2. Follow-up methods included in at least one of the visits in the 22 articles included

| Follow-up method | Frequency | Percentage |
|------------------|-----------|------------|
| ERUS             | 22        | 100%       |
| Digital examination | 20       | 90.09%     |
| CT or MRI        | 13        | 50.00%     |
| Colonography     | 11        | 50.00%     |
| Biomarkers       | 13        | 59.09%     |
| PET CT or Scintigraphy | 1       | 4.54%      |

Table 3. Percentage of LR diagnosed only by ERUS/ TRUS and local recurrence rate for the 22 articles included

| Article                        | Diagnosed only by ERUS | Local recurrence rate |
|-------------------------------|------------------------|-----------------------|
| Kwakye et al, 2019 (38)       | 71.42%                 | 11.40%                |
| Manegold et al, 2019 (16)     | NA                     | 5.70%                 |
| Li et al, 2015 (15)           | NA                     | 3.30%                 |
| Park et al, 2015 (39)         | 13%                    | 7.40%                 |
| Marken et al, 2006 (21)       | 31%                    | 11.63%                |
| Hernandez de Anda et al, 2004 (22) | 31%             | 11.63%                |
| Drudi et al, 2001 (24)        | NA                     | 47.05%                |
| Hunerbein et al, 2001 (25)    | 67%                    | 11.53%                |
| Garcia-Aguilar et al, 2000 (4) | NA                  | 24.39%                |
| Lohnert et al, 2000 (40)      | 22.00%                 | 34.30%                |
| Ross et al, 1999 (41)         | NA                     | 12.70%                |
| Novell et al, 1997 (42)       | NA                     | 15%                   |
| Rotondano et al, 1997 (43)    | 18%                    | 17.74%                |
| Hunerbein et al, 1996 (44)    | 67%                    | 17%                   |
| de Gara et al, 1995 (36)      | NA                     | 20%                   |
| Ramirez et al, 1994 (17)      | 23%                    | 20%                   |
| Romano et al, 1993 (45)       | 67%                    | 16.20%                |
| Scalpi et al, 1993 (37)       | NA                     | 9%                    |
| Dressing et al, 1990 (46)     | 33%                    | 12%                   |
| Beynon et al, 1989 (47)       | 14%                    | 25.90%                |
| Mascagni et al, 1989 (48)     | 35%                    | 14%                   |
| Hildebrandt et al, 1986 (49)  | 27%                    | 25.50%                |

| Calculated values            | 47.57 ± 12             | 15 ± 2.99             |
|                              | (25.5 to 49.7) %       | (12 to 18) %          |
Figure 2. Forrest plots graphical representation of Sensitivity and Specificity of ERUS/ TRUS (constructed with MetaDisc 1.4)

Table 4. Sensitivity and Specificity of ERUS/ TRUS calculated for the 16 articles included in meta-analysis

| Article                          | Sen. | 95% CI | Spe. | 95% CI | TP/(TP+FN) | TN/(TN+FP) |
|---------------------------------|------|--------|------|--------|------------|------------|
| Morken et al, 2006 [21]         | 0.826| 0.686 - 0.922 | 1.000| 0.794 - 1.000 | 38/46      | 16/16      |
| Drudi et al 2001 [24]           | 0.941| 0.733 - 0.999 | 1.000| 0.805 - 1.000 | 16/17      | 17/17      |
| Hönébein et al, 2001 [25]       | 0.860| 0.668 - 0.975 | 0.953| 0.842 - 0.994 | 22/25      | 41/43      |
| García-Aguilar et al, 2000 [4]  | 1.000| 0.832 - 1.000 | 1.000| 0.942 - 1.000 | 20/20      | 62/62      |
| Lohr et al 2000 [40]            | 0.793| 0.708 - 0.863 | 1.000| 0.984 - 1.000 | 92/116     | 222/222    |
| Novell et al, 1997 [42]         | 1.000| 0.839 - 1.000 | 1.000| 0.969 - 1.000 | 21/21      | 118/119    |
| Rolando et al, 1997 [43]        | 1.000| 0.715 - 1.000 | 0.980| 0.886 - 1.000 | 11/11      | 50/51      |
| Hönébein et al, 1996 [44]       | 1.000| 0.664 - 1.000 | 0.890| 0.829 - 0.934 | 9/9        | 137/154    |
| de Gara et al, 1995 [36]        | 0.714| 0.419 - 0.916 | 0.699| 0.385 - 0.803 | 10/14      | 14/23      |
| Ramirez et al, 1994 [17]        | 1.000| 0.753 - 1.000 | 0.925| 0.818 - 0.979 | 13/13      | 49/53      |
| Romano et al, 1993 [45]         | 0.667| 0.223 - 0.957 | 0.935| 0.786 - 0.992 | 4/6        | 29/31      |
| Scalpi et al, 1993 [37]         | 1.000| 0.158 - 1.000 | 0.750| 0.509 - 0.913 | 2/2        | 15/20      |
| Dressing et al, 1990 [46]       | 1.000| 0.735 - 1.000 | 0.745| 0.644 - 0.829 | 12/12      | 70/94      |
| Beynond et al, 1989 [47]        | 1.000| 0.846 - 1.000 | 1.000| 0.943 - 1.000 | 22/22      | 63/63      |
| Mascagni et al, 1989 [48]       | 0.895| 0.669 - 0.987 | 0.990| 0.946 - 1.000 | 17/19      | 100/101    |
| Hildebrandt et al, 1986 [49]    | 1.000| 0.846 - 1.000 | 1.000| 0.944 - 1.000 | 22/22      | 64/64      |

Pooled Sensitivity & Specificity: 0.883 - 0.946 - 0.987

\[ Pooled \text{Sensitivity} = 0.883, Pooled \text{Specificity} = 0.943 \]

\[ I^2 = 68.3\% \text{ for Sen. and } I^2 = 90.0\% \text{ for Spe.} \]
than 30% of instances. This result is in agreement with other systematic reviews and meta-analysis (9,19,20) which looked at ERUS/TRUS alone or in comparison with other imaging methods for preoperative staging. As of our knowledge, this is the first systematic review to focus on the effectiveness of ERUS/TRUS to diagnose LR after rectal cancer treatment. Our review has looked at the past 35 years of published literature in this area. The interest for this area changed over time, with the field starting to grow in the late 1980's, reaching a peak of research in the 1990's and a steady decrease to today. Most of the articles included in our study (10 articles) were published between 1991 and 2000 by either German or Italian research groups. In compliance with this remark, only one article had its

Table 5. Summary of positive and negative likelihood ratio of ERUS/TRUS calculated for the 16 articles included in meta-analysis

| Article                                | LR+  | 95% CI    | % Weight | LR-  | 95% CI    | % Weight |
|----------------------------------------|------|-----------|----------|------|-----------|----------|
| Morken et al, 2006 (21)                | 27.85| 1.809 - 428.77 | 4.86     | 0.186| 0.101 - 0.344 | 13.90    |
| Drudi et al 2001 (24)                  | 33.00| 2.138 - 509.33 | 4.86     | 0.086| 0.018 - 0.397 | 6.55     |
| Hunerbein et al, 2001 (25)             | 18.00| 4.851 - 73.791 | 7.08     | 0.126| 0.043 - 0.365 | 9.73     |
| Garcia-Aguilar et al 2000 (4)         | 123.00| 7.772 - 1946.5 | 4.82     | 0.024| 0.002 - 0.371 | 2.75     |
| Lohner et al 2000 (40)                 | 352.61| 22.089 - 5628.7 | 4.81     | 0.210| 0.148 - 0.298 | 16.23    |
| Novell et al, 1997 (42)                | 234.55| 14.745 - 3731.0 | 4.81     | 0.023| 0.001 - 0.354 | 2.75     |
| Rotondo et al, 1997 (43)               | 33.222| 6.833 - 161.53 | 6.68     | 0.043| 0.003 - 0.347 | 2.80     |
| Hunerbein et al, 1996 (44)             | 8.414| 5.293 - 13.377 | 8.08     | 0.056| 0.004 - 0.841 | 2.82     |
| de Gara et al, 1995 (36)               | 1.825| 0.994 - 3.353 | 7.96     | 0.469| 0.193 - 1.144 | 11.24    |
| Ramirez et al, 1994 (17)               | 11.571| 4.750 - 28.187 | 7.67     | 0.039| 0.003 - 0.593 | 2.78     |
| Romano et al, 1993 (45)                | 10.333| 2.412 - 44.272 | 6.88     | 0.356| 0.114 - 1.109 | 9.16     |
| Scalpi et al, 1993 (37)                | 3.182| 1.322 - 7.659 | 7.69     | 0.226| 0.018 - 2.872 | 3.12     |
| Dressing et al, 1990 (46)              | 3.728| 2.606 - 5.333 | 8.14     | 0.052| 0.003 - 0.767 | 2.79     |
| Beynon et al, 1989 (47)                | 125.22| 7.912 - 1981.7 | 4.82     | 0.022| 0.001 - 0.340 | 2.75     |
| Mascagni et al, 1989 (48)              | 90.368| 12.776 - 639.21 | 6.08     | 0.106| 0.029 - 0.394 | 7.89     |
| Hildebrandt et al, 1986 (49)           | 127.17| 8.035 - 2013.0 | 4.82     | 0.022| 0.001 - 0.340 | 2.75     |
| Pooled LR+                             | 21.184| 8.265 - 54.298 | 10.34    | 0.082| 0.020 - 0.420 | 1.20     |
| & I² = 91.8 % for LR+ and               |                                |          |        |        |          |
| & I² = 50.6 % for LR-                  |                                |          |        |        |          |

Figure 4. Forrest plots graphical representation Diagnostic Odds R of ERUS/TRUS (generated with MetaDisc 1.4)
protocol published in a clinical trial registry and no randomized control study was conducted to select a follow-up protocol that is safe and economic, leaving this area of interest not very accurate or with low degree of confidence. This was caused in part by the immense effort health systems put into treating patients with LR, using all available resources at hand.

As the technological advancement took place, we have remarked some patterns that are not properly expressed in numeric form. The rate of LR was higher before 2000, ranging between 20 to 30 percent, whereas after 2001, it dropped to around 10 percent. Even if the ultrasound resolution and image quality raised over the past three decades, this was not shown in the effectiveness reported by authors, with a slight drop to around 70% in the 2000’s. This might be due to the eagerness to adapt this technique in the early 1980’s and 1990’s, being less time consuming and cheaper. However, the confidence in ERUS/TRUS as a method of follow-up for LR raised in the past 20 years, with studies investigating follow-up protocols considering it a good option (21-25).

Nowadays, ultrasound probes are able to create 3D reconstructions (26,27), have higher image resolution and are available in most office settings, leading to a fast follow-up method when combined with digital examination. A recent study (28), showed that 52% of American radiation oncologists consider themselves a high utilizers of ERUS/TRUS, using it in at least 75% of patients. It is important to take into consideration that our study has looked at LR and cited literature on rectal cancer staging that mentions that ERUS/TRUS is an efficient method in assessing T-stage and not N-stage, where it was shown to not be reliable, especially after neoadjuvant treatment (19,29).

In the published literature, ERUS/TRUS...
was researched for errors and problems that might lead to wrong diagnosis and to assess the level of difficulty of this technique. The evidence in literature suggests that ERUS/TRUS is a method that has a fast learning curve, inter-operator differences are minimal, errors are manageable if discussed and it can be performed by practitioners with ranging skills and experience (30-35).

From the 28 articles that we did not include in our data analysis after the initial search, the outcomes where in favour of ERUS/TRUS as a method of rectal cancer staging or follow-up after rectal cancer, but due to our inclusion criteria they were not eligible for further study.

One of the main limitations of this study is the heterogeneity among articles included in the meta-analysis and the use of only published data available. As mentioned before, data for ERUS/TRUS use in the diagnosis of LR after RC spans over the last four decades, with technology, interests in the field and data included in the published articles greatly varying. This led to studies with 100% specificity and sensitivity, whereas other studies focusing on needle-guided biopsy emphasizing the role of this more invasive technique in favour of classical ERUS/TRUS. Another confounding factor was that some studies reported the number of procedures done and the majority focused on the number of patience and the number of LR. The studies led by de Gara et al. in 1995 (36) and Scialpi et al. in 1993 (37) are two of the main outliers in our group of articles and this might be caused by the gap that was created between the advances gained by CT and MRI in favour of US technology in the beginning of the 90's. There is a change of focus in the literature that has been highlighted by our meta-analysis, recent studies published on ERUS/TRUS used in the surveillance of LR focused on the treatment of primary rectal cancer and did not offer sufficient information for our meta-analysis. Another limitation of the current study is that authors have also looked only at Medline database and ClinicalTrials.gov registry, whereas, a future, more in-depth study should search in all available databases (BIOSIS, Scopus, WOS, Cochrane, etc.). We did not register this review in a prospective registry for systematic reviews on human studies.

We advise in favour of a multicentric randomized control trial to look for the best imaging method of follow-up for rectal cancer patients, taking into consideration patients treated with chemoradiotherapy before and/or after the surgical treatment, period of follow-up and a cost-efficiency analysis.

**Conclusion**

Based on the results of this systematic review of literature, ERUS/TRUS seems to be an efficient follow-up method for diagnosing locally recurrent rectal cancer. It is able to detect before other imaging techniques more than a third of LR, can be done in an outpatient setting and together with digital examination has a high accuracy. It is a reproducible follow-up method preferred by
practitioners, has manageable technique errors and a fast-learning curve.

**Author Contribution**

AA is the main author of this study, AA designed the research, AA and LIS collected and analysed the data. FDU contributed substantially by critical revision. AA and LIS wrote the manuscript. The manuscript has been approved by all authors.

**Conflict of Interest**

The authors have no conflict of interest to declare. None of the authors received financial support or had any institutional or other relationships that might lead to bias for this study. LIS is the recipient of a scholarship from SOF MEDICA GROUP for his educational achievements.

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