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Article (Accepted version) (Refereed)

Original citation:

DOI: 10.1016/j.jclinepi.2018.09.003

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PII: S0895-4356(18)30278-6
DOI: 10.1016/j.jclinepi.2018.09.003
Reference: JCE 9733

To appear in: Journal of Clinical Epidemiology

Received Date: 23 March 2018
Revised Date: 14 July 2018
Accepted Date: 10 September 2018


This is a PDF file of an unedited manuscript that has been accepted for publication. As a service to our customers we are providing this early version of the manuscript. The manuscript will undergo copyediting, typesetting, and review of the resulting proof before it is published in its final form. Please note that during the production process errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.
A review of NICE appraisals of pharmaceuticals 2000-2016 found variation in establishing comparative clinical effectiveness

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Key words: Single-arm Trials, Uncontrolled Trial, Drug Approval, Nonrandomised studies, Quality of evidence, National Institute of Health and Care Excellence

Conflict of Interest: DM and LO are both employed by the National Institute of Health and Care Excellence as part of the scientific advice team. There are no other conflicts of interest to declare.

Funding: This research did not receive any specific grant from funding agencies in the public, commercial, or not-for-profit sectors.

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Abstract

Objective: To identify and assess the methods for estimating comparative clinical effectiveness for novel pharmaceutical products licensed on the basis of non-RCT data and to evaluate the corresponding NICE recommendations.

Methods: Our identification strategy was two-fold. First, we reviewed all NICE appraisals between 2010 and 2016 and identified technologies where comparative clinical effectiveness estimates were calculated using non-RCT data. Second, we checked if NICE appraisals completed from 2000 to 2010 had included pharmaceuticals that were granted EMA marketing authorisation without RCT data between 1999-2014. Information was extracted on the methods used as well as the corresponding NICE recommendations. We also collected information on the rationale for utilising non-RCT data in NICE appraisals.

Results: Of 489 individual pharmaceutical technologies assessed by NICE, 22 (4%) used non-RCT data to estimate comparative clinical effectiveness. Methods for establishing external controls in such studies varied: 13 (59%) used published trials, 6 (27%) used observational data, 2 (9%) used expert opinion, and 1 (5%) used a responder vs non-responder analysis. Only 5 (23%) used a regression model to adjust for covariates. We did not observe a notable difference in the proportion of pharmaceutical technologies that received a positive recommendation from NICE whether the decision was based on RCT or non-RCT data. (83% vs 86%)

Conclusions: To date, the small number of appraisals by NICE based on non-RCT data did not result in substantially different treatment decisions. The majority of the technologies appraised on the basis of non-RCT data either received a positive recommendation or a positive recommendation with restrictions. The methods used to calculate comparative clinical effectiveness estimates varied, highlighting the need to establish clear guidance.
What is new?

Key findings

- Between 2000-2016, 22 of the 489 (4%) individual pharmaceutical technologies assessed by NICE were based upon comparative clinical effectiveness estimates calculated using non-RCT data. Out of these:
  - 11 (50%) were included in technology appraisals published in either 2015 or 2016.
  - 10 (45%) received a positive recommendation from NICE, 9 (41%) received an optimised recommendation and a further 3 (14%) received a negative recommendation.
  - 14 (64%) appraisals calculated comparative clinical effectiveness estimates using a naive unadjusted indirect comparison against an external control, not utilising any regression methods. This may reflect the limited availability of individual patient level data to adjust for covariates.

What this adds to what was known?

- Despite non-RCT data leading to significant uncertainty in quantifying clinical benefit, we only observed a small differences in the proportion of pharmaceutical technologies that received a positive or negative recommendation from NICE when comparing decisions based on RCT or non-RCT data.

What is the implication and what should change now?

- Clear guidance is needed to establish the comparative clinical effectiveness of pharmaceuticals with non-RCT data.
- There is a need to monitor and follow-up the real-world comparative clinical and cost effectiveness of pharmaceutical technologies recommended on the basis of non-RCT data.
1. Introduction

The gold standard for establishing the clinical effectiveness and safety of a pharmaceutical technology is to conduct a randomised controlled trial (RCT). RCTs are the mainstay of research and development of new medicines, and are used to establish reliable comparative efficacy estimates.\(^{(1)}\) The random allocation of patients between an intervention and control arm reduces confounding as compared to other types of study design where there is no randomisation.\(^{(2,3)}\) This is reflected in the National Institute for Health and Care Excellence (NICE) guidance for technology appraisals which states that RCTs are “considered to be most appropriate for measures of relative treatment effect” and outlines that the “problems of confounding, lack of blinding, incomplete follow-up and lack of a clear denominator and endpoint occur more commonly in non-randomised studies and non-controlled trials than in RCTs”.\(^{(4)}\)

Clinical equipoise, defined as the absence of certainty about the superiority of alternative treatment options for a given indication, is a fundamental ethical criterion to randomly allocate participants to different treatment groups.\(^{(5)}\) However, this criterion may not be necessary in the context of precision medicine with targeted therapies, where earlier studies have demonstrated a large magnitude of treatment effect. In such cases, some observers have postulated that RCTs may not be necessary.\(^{(6–8)}\) Additionally, in the context of small populations, RCTs may be unethical and misleading, as results could be statistically underpowered.\(^{(9)}\) As a result, drug licensing agencies such as the European Medicines Agency (EMA) may grant marketing authorisations to pharmaceutical technologies with no RCT data, when there is certainty that the product’s benefit outweighs potential harm.\(^{(10)}\)

Once products are on the market, health technology assessment (HTA) agencies such as the National Institute for Health and Care Excellence (NICE) are responsible for evaluating not only the relative benefits but also the cost-effectiveness of novel technologies compared to existing alternatives in established use in clinical practice. Reliance on non-RCT data poses significant challenges for HTA agencies that are faced with increased uncertainty regarding the comparative clinical effectiveness, and therefore the cost-effectiveness estimates of pharmaceutical products.\(^{(11,12)}\)

In the absence of a controlled trial, alternative methods are required to generate relative clinical effectiveness estimates against comparator agents. Such estimates are an essential component for economic evaluation analyses conducted to establish the value of new agents. For example, an external control could be considered in an economic evaluation model using either historical data or a self-control.\(^{(13)}\) According to technical guidance developed by NICE’s Decision Support Unit, use of a regression model is the preferred option to adjust for the effects of covariates when using an external control, although this is only possible when individual patient-level data are available for both the non-controlled trial and external control.\(^{(14)}\)

Our objective in this paper was to review NICE appraisals between 2000 and 2016 that calculated comparative clinical effectiveness estimates using non-RCT data. We reviewed the methods used and compared the final published committee recommendation for these technologies versus those where comparative clinical effectiveness estimates were calculated using RCT data.
2. Methods

2.1. Identification of Pharmaceutical Technologies
We adopted a two-pronged approach to identify pharmaceutical technologies whereby comparative clinical effectiveness estimates in NICE technology appraisals were calculated using non-RCT data. First, one researcher systematically reviewed all publicly available guidance documents from NICE single technology appraisal (STA), multiple technology appraisal (MTA) and highly specialised technology (HST) processes published between January 2010 and December 2016. Second, we reviewed all NICE STA, MTA, HST guidance published between January 2000 and December 2016 which appraised pharmaceutical technologies listed within two previously published systematic reviews of pharmaceutical technologies which were granted EMA marketing authorisation without RCT data between 1999-2014.(10,15)

For each technology appraisal, the corresponding clinical and economic evidence within NICE committee papers and evidence review group (ERG) reports were reviewed. NICE committee members are selected from the Institute itself, the National Health Service (NHS), patient and carer organisations, academia and the pharmaceutical industry.(16) The ERG reports are produced by a group of independent experts in academia, and commissioned by the National Institute for Health Research (NIHR) to review and critique both the clinical and cost-effectiveness evidence available for each technology under appraisal.(4)

As the NICE MTA process considers multiple pharmaceutical technologies for the same indication, each individual pharmaceutical technology under MTA with its own corresponding clinical and economic evidence base was reviewed separately.

We excluded all non-pharmaceutical technology appraisals and those which were subsequently updated or terminated due to manufacturer non-submission.

2.2. Eligibility Criteria
Pharmaceutical interventions reviewed by the NICE STA, MTA and HST appraisal processes were screened and categorised by one researcher as either a non-RCT or an RCT-based technology. A non-RCT based technology was defined as a pharmaceutical technology whereby the comparative clinical effectiveness estimates used within the economic evaluation model were calculated using non-RCT data. Non-RCT data could either be obtained from uncontrolled studies (i.e., a single-arm trial or a trial without a concurrent comparator group) or the intervention arm of an RCT interpreted as a single-arm trial by NICE. The latter occurred when the original comparator included in the RCT was deemed not to be relevant to the NICE decision scope (as specified in the published appraisal report).

We relied on the ERG's final economic evaluation model. The ERG may choose to alter the economic evaluation model submitted by the manufacturer to correct errors, consider different clinical or cost inputs, or modify the structure of the model. The ERG's final economic evaluation model is therefore the most comprehensive reflection of available evidence which ultimately informs the NICE assessment.

The final sample of technologies was checked and confirmed by a senior member of the research team.

2.3. Data Extraction
One researcher reviewed the appraisal guidance and ERG reports and collected data on (1) the methods for establishing comparative clinical effectiveness estimates, (2) NICE recommendations, and (3) NICE committee comments.
2.3.1. Methods to establish comparative clinical effectiveness estimates
For each technology in our sample, within the final ERG’s economic evaluation, an unanchored indirect comparison was utilised to estimate comparative clinical effectiveness. The external control and methods used to estimate comparative clinical effectiveness in the economic model was identified. It was also determined if a meta-analysis or regression model was used in the final economic model to adjust for covariates.

2.3.2. NICE Recommendations
One researcher reviewed the NICE committee papers to characterise the corresponding recommendations for all technologies, including both the committee decision outcome and any use of patient access schemes. The committee decision outcome could be ‘Recommended’ (approved with no restrictions), ‘Optimised’ (approved within a specified patient subgroup), ‘Not Recommended’ (not approved for routine use in the NHS), or ‘Only in Research’ (approved within a research setting). Many recommendations by NICE are based on a patient access scheme in which the cost-effectiveness of a technology under appraisal is improved by offering the technology at a discounted price. The NICE recommendations were reviewed for all technology appraisals published by NICE between January 2000 and December 2016.

2.3.3. NICE Committee Comments
For all technologies in our sample, one researcher reviewed the key conclusions of NICE committee documents to highlight any additional factors considered as well as concerns regarding the clinical evidence base and/or incremental cost-effectiveness ratio (ICER) estimates to gain an insight into the committee’s decision-making process.

3. Results

3.1. Identified Pharmaceutical Technologies
We identified a total of 429 NICE technology appraisals between January 2000 and December 2016 (Figure 1). Of these, we excluded 120 as they either included non-pharmaceutical technologies (37), were subsequently updated (57) or were terminated due to manufacturer non-submission (26). In the latter two cases, NICE committee papers were not publicly available. Reviewing the EMA initial marketing authorisation documents revealed 4/26 (15%) of these terminated technologies were approved on the basis of a pivotal trial with non-RCT data. All 26 terminated technologies were due to manufacturer non-submission.

The remaining 309 technology appraisals included a total of 489 individual pharmaceutical products. Of these 489 pharmaceutical technologies, 22 (4%) individual pharmaceuticals across 20 technology appraisals were based on non-RCT data and therefore met our eligibility criteria. 12 (55%) had an oncology indication, 6 (27%) had a hepatology indication, 3 (14%) had a rheumatology indication and 1 (5%) had an immunology indication.

3.2. Methods used to establish Comparative Efficacy
When choosing an external control to establish comparative clinical effectiveness estimates in the economic model, 13 (59%) used previously published trials, 6 (27%) used observational data, 2 (9%) used expert opinion and 1 (5%) used a responder vs non-responder analysis. (Table 2). Only 5 (23%) appraisals utilised a regression model, and only 6 (27%) appraisals used a meta-analysis to combine results from multiple studies. Comparative clinical effectiveness estimates for 14 pharmaceutical technologies (64%) were
calculated using a naive unadjusted indirect comparison against an external control, possibly due to a lack of individual patient level data.

### 3.3. NICE Recommendations

There was small differences between technologies in our sample (appraised on the basis of non-RCT data) vs technologies with RCT data receiving the NICE Committee decision outcome of ‘Recommended’ 10/22 vs 289/467 (45% vs 62%) or ‘Not Recommended’ 3/22 vs 81/467 (14% vs 17%) (Table 1). Technologies in our sample were more than twice as likely to receive the NICE Committee decision outcome of ‘Optimised’ as compared to those with RCT data (9/22 vs 91/467; 41% vs 19%). The ‘Only In Research’ designation was not used for non-RCT-based pharmaceutical technologies and for 6/467 (1%) of RCT-based pharmaceutical technologies.

Technologies with non-RCT data were more likely to utilise a patient access scheme as compared to those with RCT data; 7/22 vs 111/467 (32% vs 24%) (Table 1). All patient access schemes were financially-based except for one for a non-RCT-based technology. This scheme relied on a combination of a financial and performance-based patient access scheme, i.e. a managed access agreement.(17)

### 3.4. NICE Committee Comments

Several factors were considered by NICE Committees when evaluating technologies with non-RCT data. The most frequent factors were significant unmet need (11/22, 50%), a small patient population (6/22, 27%) and cases when early trials had shown substantial benefit (2/22, 9%). The small sample size limited the possibility to explore the association between NICE committee recommendations and these factors (Table 1) All committees explicitly highlighted concerns regarding the clinical evidence for all 22 technologies in our sample. These concerns included the immaturity of data, and the uncertainty associated with the lack of a direct comparator. Conversely, issues regarding the cost-effectiveness of each technology were more inconsistent, they were present for the 2 ‘Not Recommended’ technologies where an ICER estimate was available (100%, 2/2), the majority of the 9 ‘Optimised’ technologies (78%, 7/9) and rarely present for the 10 ‘Recommended’ technologies (20%, 210). (Table 1) Concerns were typically raised when the associated ICER estimate was above an acceptable threshold.

### 4. Discussion

#### 4.1. Summary of Findings

Our review of NICE appraisals conducted between 2000 and 2016 identified 22 pharmaceutical technologies that relied on non-RCT data when generating comparative efficacy estimates used in the economic model. In these instances, we did not identify a consistent methodological approach to compare the technology to its comparators specified in the decision scope. Whilst existing guidelines recommend the use of statistical methods to adjust for covariates,(14,18) a regression model was used in only 5 out of 22 appraisals identified in our review. Final recommendations did not differ for technologies with and without RCT data.

Previous research has indicated that the strongest predictor for a NICE recommendation is the ICER estimate.(19) Other factors which may influence NICE recommendations include severity of underlying illness, end-of-life considerations, disadvantaged populations, unmet need and paediatric indications.(20) Our findings are consistent with the previous literature. We found no notable differences between technologies appraised on the basis of non-RCT or RCT data receiving a positive (86% vs 83%) or negative (14% vs 17%) recommendation from NICE. We found that NICE Committees considered several additional factors when
appraising technologies on the basis of non-RCT data, including unmet clinical need, small patient populations, and large treatment effects. Amongst the positive recommendations there was a higher proportion of technologies with non-RCT data receiving the ‘Optimised’ decision outcome (41% vs 19%), likely reflecting the fragmented nature of the target patient population. In these circumstances, a patient population may be defined by a disease stage or other subgroup and be limited in numbers; conducting an RCT may therefore be challenging.

Reviewing a recent EMA report, it was found that 6/22 (27%) of these technologies were granted conditional marketing authorisation by the EMA.\(^{(21)}\) Despite several alternative strategies to mitigate the uncertainty associated with relative clinical effectiveness estimates derived from non-RCT data, they were seldom used by manufacturers or ERGs. Firstly, the ‘Only In Research’ recommendation was not used for any non-RCT based technologies. Although whilst the ‘Only In Research’ designation may be appropriate in some cases, this must be balanced against delayed access to medicines for patients.\(^{(22)}\) Secondly, patient access schemes offer another option to mitigate uncertainty regarding value of new technologies. Interestingly, there was only a small difference between technologies with and without RCT data utilising a patient access scheme. Furthermore, only one of these schemes was performance-based. This may reflect the methodological challenges associated with collecting real-world data following market entry.\(^{(23)}\)

### 4.2. Implications

To combat significant uncertainty associated with approving technologies based on non-RCT data alone, novel risk sharing approaches are needed. For example, NICE, NHS England, and the UK Department of Health have launched a new Cancer Drugs Fund (CDF), which fund oncology medications while further data are collected. For drugs recommended within the CDF, a managed access agreement (MAA) is agreed with both a data collection and a commercial agreement.\(^{(24)}\) Similar initiatives to address clinical uncertainty are also being considered for non-oncology medications. Furthermore, the ADAPT SMART project has begun to explore how adaptive pathways and performance based managed access schemes could be utilised.\(^{(25)}\)

Although only a small number of pharmaceutical technologies were appraised on the basis of non-RCT data during our study period, the majority were in the last few years. Recent research has shown that larger effect sizes in noncontrolled studies are associated with higher rates of EMA licensing approval.\(^{7}\) Whilst this offers useful insight, clear criteria are needed to specify the circumstances in which noncontrolled trials are appropriate. Currently, the uncertainty in guidance represents a significant challenge for the pharmaceutical industry, regulatory authorities and HTA organisations. A collaborative technical advisory group involving both regulatory authorities such as the EMA and HTA organisations could begin efforts towards outlining guidance for a more consistent approach to both assessing the suitability of non-RCT data in evidence submissions and accruing further evidence over time.

### 4.3. Limitations

This study has a number of limitations. First, although our review comprehensively captures NICE appraisals of technologies that received EMA approvals without RCT data, we may have missed technologies from 2000 to 2009 if NICE considered the comparator arm of some trials to be irrelevant for the decision scope during this period. Second, while one researcher was involved in screening the appraisals extracting data from technology appraisals and associated documents, a senior member of the research team verified the sample. Third, our sample size was small, as technologies appraised on the basis of RCT data constitute the majority of NICE appraisals. However, half of the technologies without RCT data in our sample were appraised in the last 2 years.
5. Conclusion

From 2000 to 2016, 22 technologies were appraised by NICE based on non-RCT data. The methods used to calculate comparative efficacy estimates in the absence of comparative trials varied. While regression methods can decrease the uncertainty associated with the evidence base of these technologies, this relies upon the availability of individual patient-level data for any external control selected. Only a minority of the technologies appraised on the basis of non-RCT data received a negative recommendation.

Authors’ Contributions: The conception and design of the study were completed by MA, HN, DM, LO and EM. Acquisition of data was done by MA and HN. Analysis of data was done by MA. Interpretation of data was handled by MA and HN. Drafting of manuscript was completed by MA and HN. Critical revisions were done by all authors. All authors gave final approval of the completed manuscript.

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Table 1. Characteristics of technology appraisals with economic evaluations using non-RCT data to establish comparative clinical efficacy

<p>| No  | Year | NICE Programme | Technology | Indication                                                                 | Therapeutic Area | PAS | Additional Factors considered by NICE Committees | NICE Committee Concerns | Efficacy Estimate | ICER Estimate |
|-----|------|----------------|------------|----------------------------------------------------------------------------|------------------|-----|-----------------------------------------------|--------------------------|------------------|---------------|----------------|
| 178 | 2009 | MTA            | Sunitinib  | Advanced and/or metastatic renal cell carcinoma (2nd line)                 | Oncology         | Yes | Significant Unmet Need                       | Yes                      | N/A              |               |                |
| 202 | 2010 | STA            | Ofatumumab | Chronic lymphocytic leukaemia refractory to fludarabine and alemtuzumab    | Oncology         | Yes | Early trial showed substantial benefit       | Yes                      | Yes              |               |                |
| 209 | 2010 | MTA            | Imatinib   | Unresectable and/or metastatic gastrointestinal stromal tumours            | Oncology         | No  | Small Patient Population                     | Yes                      | Yes              |               |                |
| 195 | 2010 | MTA            | Adalimumab | Rheumatoid arthritis after the failure of a TNF inhibitor                  | Rheumatology     | No  | Nil                                           | Yes                      | Yes              |               |                |
| 195 | 2010 | MTA            | Etanercept | Rheumatoid arthritis after the failure of a TNF inhibitor                  | Rheumatology     | No  | Nil                                           | Yes                      | Yes              |               |                |
| 195 | 2010 | MTA            | Infliximab | Rheumatoid arthritis after the failure of a TNF inhibitor                  | Rheumatology     | No  | Nil                                           | Yes                      | Yes              |               |                |
| 330 | 2015 | STA            | Sofosbuvir | Chronic hepatitis C                                                        | Hepatology       | No  | Early trial showed substantial benefit       | Yes                      | Yes              |               |                |
| 363 | 2015 | STA            | Ledipasvir | Chronic hepatitis C                                                        | Hepatology       | No  | Significant unmet need                       | Yes                      | Yes              |               |                |
| 364 | 2015 | STA            | Daclatasvir| Chronic hepatitis C                                                        | Hepatology       | No  | Significant unmet need                       | Yes                      | Yes              |               |                |
| HST1| 2015 | HST            | Eculizumab | Atypical haemolytic uraemic syndrome                                       | Nephrology       | No  | Small patient population                      | Yes                      | Yes              |               |                |
| 408 | 2016 | STA            | Pegaspargase| Acute lymphoblastic leukaemia                                              | Oncology         | No  | Paediatrics: Nil Adults: Small Population     | Yes                      | No               |               |                |
| 410 | 2016 | STA            | Tamlimogene | Unresectable metastatic melanoma                                           | Oncology         | Yes | Small patient population. Significant unmet need | Yes                      | No               |               |                |
| 23  | 2001 | MTA            | Temozolomide| Recurrent malignant glioma                                                 | Oncology         | No  | Small patient population                      | Yes                      | Yes              |               |                |
| 86  | 2004 | MTA            | Imatinib   | Unresectable and/or metastatic gastrointestinal stromal tumours            | Oncology         | No  | Significant Unmet Need                       | Yes                      | No               |               |                |</p>
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<td>Oncology</td>
<td>Yes</td>
<td>Small patient population. Significant Unmet Need</td>
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<td>Yes</td>
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**PAS:** Patient Access Scheme  **ERG:** Evidence Review Group  **NICE:** National Institute for Health and Care Excellence  **ICER:** Incremental Cost-Effectiveness Ratio  **STA:** Single Technology Appraisal  **MTA:** Multiple Technology Appraisal  **HST:** Highly-Specialised Technologies.
Table 2. Methods used for technology appraisals with economic evaluations using non-RCT data to establish comparative clinical efficacy

<p>| No | Year | Name | Intervention Efficacy Data | Comparators Efficacy Data | Comparison Method | Meta-Analysis | Regression Model |
|----|------|------|-----------------------------|---------------------------|-------------------|---------------|-----------------|-----------------|
| 178 | 2009 | Sunitinib | Two Uncontrolled Studies | No Comparison made | No Comparison made | No | No |
| 202 | 2010 | Ofatumumab | Single uncontrolled study* (Responders) | Single uncontrolled study* (Non-responders) | Responder vs Non-Responder Analysis | No | Yes |
| 209 | 2010 | Imatinib 600mg/800mg | Multiple uncontrolled studies | Observational Study | Naïve unadjusted Indirect comparison | No | No |
| 195 | 2010 | Adalimumab | Single uncontrolled study | Multiple uncontrolled trials and RCTs | Naïve unadjusted Indirect comparison | Yes | No |
| 195 | 2010 | Etanercept | Single uncontrolled study | Multiple uncontrolled trials and RCTs | Naïve unadjusted Indirect comparison | Yes | No |
| 195 | 2010 | Infliximab | Single uncontrolled study | Multiple uncontrolled trials and RCTs | Naïve unadjusted Indirect comparison | Yes | No |
| 330 | 2015 | Sofosbuvir | Multiple uncontrolled studies | Multiple uncontrolled studies and RCTs | Naïve unadjusted Indirect comparison | No | No |
| 363 | 2015 | Ledipasvir-Sofosbuvir | Multiple uncontrolled studies | Multiple uncontrolled studies and RCTs | Naïve unadjusted Indirect comparison | No | No |
| 364 | 2015 | Daclatasvir | Multiple uncontrolled studies | Multiple uncontrolled studies and RCTs | Naïve unadjusted Indirect comparison | No | No |
| HS T1 | 2015 | Eculizumab | Single uncontrolled study | Observational dataset | Naïve unadjusted Indirect comparison | No | No |
| 408 | 2016 | Pegaspargase | Paediatric: One uncontrolled study Adult: Multiple uncontrolled studies | Paediatric: Multiple single-arms of RCTs Adult: Expert Opinion | Paediatrics: Naïve unadjusted Indirect comparison Adult: Expert Opinion ≤25 years-Yes (Comparators) &gt;25 years-No | No | No |
| 410 | 2016 | Tamlimogene Laherparepvec | Single-arm of an RCT | Multiple RCTs | Adjusted Indirect Comparison | No | Yes |
| 23 | 2001 | Temozolomide | Single-arm of an RCT | Multiple uncontrolled studies | Naïve unadjusted indirect comparison | Yes | No |
| 86 | 2004 | Imatinib | Single uncontrolled study | Observational Study | Naïve unadjusted Indirect comparison | No | No |
| 185 | 2010 | Intravenous Trabectin | Single uncontrolled study | Observational dataset | Adjusted Indirect Comparison | No | Yes |
| 246 | 2012 | Pharmalgen | Multiple uncontrolled studies | Observational Study | Naïve unadjusted Indirect comparison | Yes | No |
| 300 | 2013 | Peginterferon alfa | Single uncontrolled study | Expert Opinion | Expert Opinion | No | No |</p>
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</tr>
<tr>
<td>401</td>
<td>2016</td>
<td>Bosutinib</td>
<td>Single uncontrolled study</td>
<td>Observational Study</td>
<td>Naïve unadjusted Indirect comparison</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>416</td>
<td>2016</td>
<td>Osimertinib</td>
<td>Multiple uncontrolled studies</td>
<td>One RCT</td>
<td>Adjusted indirect comparison</td>
<td>Yes</td>
<td>Yes</td>
</tr>
</tbody>
</table>

*This was the same study

**NICE:** National Institute for Health and Care Excellence  **RCT:** Randomised Controlled Trial
Table 3. NICE Committee Recommendations

<table>
<thead>
<tr>
<th>Decision Outcome</th>
<th>Non-RCT based Technologies</th>
<th>RCT based Technologies</th>
</tr>
</thead>
<tbody>
<tr>
<td>Recommended</td>
<td>10/22 (45%)</td>
<td>289/467 (62%)</td>
</tr>
<tr>
<td>Not Recommended</td>
<td>3/22 (14%)</td>
<td>81/467 (17%)</td>
</tr>
<tr>
<td>Optimised</td>
<td>9/22 (41%)</td>
<td>91/467 (19%)</td>
</tr>
<tr>
<td>Only in Research</td>
<td>0/22 (0%)</td>
<td>6/467 (1%)</td>
</tr>
</tbody>
</table>

**Patient Access Scheme**

| Patient Access Scheme   | 7/22 (32%)                 | 111/467 (24%)          |

**RCT**: randomised controlled trial.
Figure 1. Flow Diagram of Study Selection Process for Appraisals in the Review

Between 2000-2016 NICE published 429 Technology Appraisals
- STA: 276
- MTA: 150
- HST: 3

Excluded 120:
- 37 Devices/Non-Pharmaceutical
- 57 Updated/Replaced
- 26 Terminated-Non Submission

Remaining 309 Technology Appraisals
Covering a total of:
- 489 Individual Pharmaceutical Technologies

A review of clinical and cost-effectiveness sections within Evidence Review Group and NICE Committee Papers of all technology appraisals for pharmaceutical technologies contained within:


Results
- 22 (4.5%) Pharmaceutical Technologies in which comparative clinical effectiveness is established with uncontrolled data
- 467 (95.5%) Pharmaceutical Technologies in which comparative clinical effectiveness is established with RCT data

NICE: National Institute for Health and Care Excellence; STA: single-technology appraisals; MTA: multiple-technology appraisals; HST: highly-specialised technologies;