A Comprehensive Algorithm for Approval of Health Technologies With, Without, or Only in Research: The Key Principles for Informing Coverage Decisions

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ABSTRACT

Background: The value of evidence about the performance of a technology and the value of access to a technology are central to policy decisions regarding coverage with, without, or only in research and managed entry (or risk-sharing) agreements. Objectives: We aim to outline the key principles of what assessments are needed to inform “only in research” (OIR) or “approval with research” (AWR) recommendations, in addition to approval or rejection. Methods: We developed a comprehensive algorithm to inform the sequence of assessments and judgments that lead to different types of guidance: OIR, AWR, Approve, or Reject. This algorithm identifies the order in which assessments might be made, how similar guidance might be arrived at through different combinations of considerations, and when guidance might change. Results: The key principles are whether the technology is expected to be cost-effective; whether the technology has significant irreversible costs; whether additional research is needed; whether research is possible with approval and whether there are opportunity costs that once committed by approval cannot be recovered; and whether there are effective price reductions. Determining expected cost-effectiveness is only a first step. In addition to AWR for technologies expected to be cost-effective and OIR for those not expected to be cost-effective, there are other important circumstances when OIR should be considered. Conclusions: These principles demonstrate that cost-effectiveness is a necessary but not sufficient condition for approval. Even when research is possible with approval, OIR may be appropriate when a technology is expected to be cost-effective due to significant irreversible costs.

Keywords: cost-effectiveness, coverage with evidence development, health technology assessment, only in research.

Introduction

The value of evidence about the performance of a technology and the value of access to a technology are central to policy decisions regarding coverage with, without, or only in research and managed entry (or risk-sharing) agreements. Decisions about health technologies are increasingly being made close to license when the evidence base to support the technology is least mature. This is partly linked to changes in the regulatory landscape in which regulators and payers have produced new approaches in an effort to provide patients with timely access to new medicines. For example, the European Medicines Agency has introduced the “Adaptive Pathways” approach that is open to interventions in the early stages of development [1,2]. Inevitably, this means that coverage decisions in many different jurisdictions and health care systems are being made when there may be substantial uncertainty surrounding the technology’s effectiveness, cost-effectiveness, and potential for harm. In these circumstances, further evidence may be particularly valuable because it can lead to better decisions that improve patient outcomes and/or reduce resource costs. Establishing the key principles for “only in research” (OIR), defined as when a technology is approved only within the context of a suitable research study, and “approval with research” (AWR), which refers to approval while research is being conducted, will enable coverage decisions to be addressed in an explicit and transparent manner. This has wide relevance to different types of health care systems, helping to inform the questions posed by coverage with evidence development and managed entry agreements [3–6].

If the resources available for health care are limited, then approving a more costly technology will displace other activities that would have otherwise generated improvements in health for other patients [7]. If the objective of a health care system is to improve health outcomes across the population it serves, then
even if a technology is expected to be more effective, the health gained must be compared with the health expected to be forgone elsewhere as a consequence of additional costs; that is, a cost-effective technology will offer positive net health effects (NHEs) [8–10]. An assessment of expected cost-effectiveness or NHEs relies on evidence about effectiveness, impact on long-term overall health and potential harms, as well as additional health care costs together with some assessment of what health is likely to be forgone as a consequence (the cost-effectiveness threshold) [11]. In health systems without administrative budgets, there are often constraints on the growth in health care expenditure. Even where there are no constraints, the same principles are likely to apply but the opportunity costs may manifest in terms of non-health expenditure, for example, through increased insurance premiums, co-payments, or taxation.

Such assessments are inevitably uncertain and without sufficient and good quality evidence, subsequent decisions about the use of technologies will also be uncertain. There will be a chance that the resources committed by the approval of a new technology may be wasted if the expected positive NHEs are not realized. Equally, rejecting a new technology will risk failing to provide access to a valuable intervention if the NHEs prove to be greater than expected. Therefore, if the social objective is to improve the overall health for both current and future patients, then the need for and value of additional evidence is an important consideration when making decisions about the use of technologies [12–14]. This is even more critical once it is recognized that the approval of a technology for widespread use might reduce the prospects of conducting the type of research that would provide the evidence needed [15]. In these circumstances, there will be a trade-off between the NHEs to current patients from early access to a cost-effective technology and the health benefits to future patients from withholding approval until valuable research has been conducted [16].

Research also consumes valuable resources that could have been devoted to patient care or to other more valuable research priorities. Uncertain events in the near or distant future may also change the value of the technology and the need for evidence [17]. In addition, implementing a decision to approve a new technology may commit resources that cannot subsequently be recovered if guidance changes in the future [18–22]. Guidance about the use of health technologies will depend on whether the benefits of research are likely to exceed the costs and whether any benefits of early approval are greater than withholding approval until additional research is conducted or other sources of uncertainty are resolved.

The purpose of this article was to outline the key principles of what assessments are needed to inform coverage decisions. The starting point is an assessment of expected cost-effectiveness from an underlying extra-welfarist approach, which identifies improvements in health as an important objective of health care provision [23]. This is the approach that has been adopted by major decision-making bodies such as the National Institute for Health and Care Excellence in the United Kingdom, the Pharmaceutical Benefits Advisory Committee in Australia, and the Canadian Agency for Drugs and Technologies in Health. These principles do not presuppose how the assessments might be informed in terms of the methods of analysis or how different aspects of health gained and forgone might be measured and valued. This distinction between principles and methods of analysis is important because different health care systems (or decision-making bodies) are likely to vary in terms of the social values they apply as well as the time and resources available to carry out an appraisal of a health technology and may adopt different methods of analysis to inform the same question. On the basis of these principles, we present a comprehensive algorithm that demonstrates how the sequence of assessments and judgments can lead to different types of guidance: OIR, AWR, Approve (without research), or Reject (without research). Full details on the development of this comprehensive framework and algorithm have been described elsewhere [24], and we refer the reader to this fuller description for a deeper insight. An illustration of how the assessments can be informed in terms of the methods of analysis is also presented elsewhere, with application to a number of case studies [24,25].

### Key Principles and Assessments Needed

The key principles fall into a number of broad areas: 1) whether the technology is expected to be cost-effective; 2) whether the technology has significant irrecoverable costs; 3) whether additional evidence/research is needed; 4) whether research is possible with approval and whether there are significant (opportunity) costs that once committed by approval cannot be recovered; and 5) whether there are any effective price reductions offered. These key principles can be represented by a sequence of assessments and judgments, which are summarized as a seven-point checklist [24,25]:

1. Is the technology expected to be cost-effective?
2. Are there significant irrecoverable costs?
3. Does more research seem worthwhile?
4. Is the research possible with (without) approval?
5. Will other sources of uncertainty resolve over time?
6. Are the benefits of research greater than the costs?
7. Are the benefits of approval greater than the opportunity costs?

Based on estimates of expected NHE at each point of this checklist, a judgment Yes or No can be made. For example, a judgment at point 1 uses standard cost-effectiveness analysis to estimate the NHE, a judgment at point 2 assesses the impact of irrecoverable costs on NHE, and a judgment at points 3 and 4 uses probabilistic sensitivity analysis and methods of value of information analysis (the reader is referred to McKenna et al. [25] for the methods of analysis for each point of the checklist). Guidance will depend on the combined effect of these assessments. In some cases, all seven assessments may not be necessary as earlier decisions lead directly to guidance.

Figures 1 to 3 show the sequence of assessments and decisions that lead to a particular category and type of guidance, represented as a comprehensive algorithm. Four broad categories of guidance are represented within the algorithm: Approve, AWR, OIR, and Reject. Each different type of guidance illustrates how similar guidance might be arrived at in different ways, helping to identify the particular combination of considerations that underpin the guidance. Figure 1 is for technologies without significant irrecoverable costs, whereas Figures 2 and 3 are for technologies with significant irrecoverable costs. Figures 2 and 3 are separated further on the basis of whether the technology is expected to be cost-effective and research is needed (Fig. 2) or whether it is not expected to be cost-effective and research may or may not be needed (Fig. 3). The key principles underpinning each of these assessments are now described in the sections that follow. The reader is referred to Claxton et al. [24] for a more detailed explanation.

### Technologies Without Significant Irrecoverable Costs

Although some element of cost which once committed cannot be subsequently recovered is almost always present in the evaluation of technologies, the sequence of assessments and decisions for these technologies is relatively straightforward compared
with other technologies judged to have “significant” irrecoverable costs associated with approval. Significant irrecoverable costs depend on the commitment of upfront costs and whether there is sufficient flexibility in when a patient can initiate treatment (e.g., if treatment can be delayed until uncertainty is resolved, then the commitment of these irrecoverable costs can be avoided). Technologies without significant irrecoverable costs are considered first as the simpler case (see Fig. 1).

Technologies expected to be cost-effective

The sequence of assessment is assumed to start with cost-effectiveness and expected incremental NHE for the technology relative to its comparators because this is the starting point used by many decision-making bodies such as the National Institute for Health and Care Excellence, the Pharmaceutical Benefits Advisory Committee, and the Canadian Agency for Drugs and Technologies in Health. To avoid binary decisions of Approve or Reject under conditions of uncertainty, the need for additional evidence should be assessed. An assessment is also required of whether the research needed to provide this evidence is possible if the technology is approved for widespread use while the research is being conducted.

Need for evidence. Some initial assessment about whether further research might be potentially worthwhile is important because a “No” at this point can avoid further assessments; for example, a technology offering substantial and well-evidenced health benefits at modest additional cost is likely to exhibit little uncertainty about whether the expected population NHE is positive. In these circumstances, further research may not be worthwhile, so guidance to approve could be issued (e.g., Approve⁴ in Fig. 1). If additional evidence is needed and further research might be worthwhile, then further assessments and decisions are required before guidance can be issued. Critically, some assessment is required of the type of evidence that is needed and whether or not the type of research required to provide it is likely to be conducted if approval is granted [24,26].

Where research is possible with approval. If research is possible with approval, some assessment of the long-term benefits of research is required, including 1) the likelihood that the type of research needed will be conducted; 2) how long the results of research will take to report; and 3) how much of the uncertainty might be resolved by the research [15]. An assessment of other sources of uncertainty that will resolve over time is also needed, for example, changes in prices or the launch of new technologies [17]. These sources of uncertainty will influence the future benefits of research. For example, if the price of the technology is likely to fall significantly before or shortly after research reports, then the benefits, once the research reports, might be very limited. In these circumstances, it might be better to approve (rather than AWR) and reconsider whether and what type of research is needed at a later date once these uncertainties have resolved. The judgment of whether the long-term benefits of research are likely to exceed its expected costs determines guidance, with AWR₁ and Approve₁ in Figure 1 dependent on “Yes” and “No,” respectively.

Where research is not possible with approval. The type of research needed may not be possible once a technology is
approved for widespread use; for example, randomized clinical trials may not be possible due to ethical concerns, recruitment problems, and limited incentives for manufacturers. In these circumstances, the expected benefits of approval to current patients must be balanced against the benefits to future patients from withholding approval until the research is conducted. The same assessments of the long-term value of research and the impact of other sources of uncertainty are still required. If the benefits of research are judged to be less than the costs (i.e., research is not worthwhile anyway), the technology can be approved on the basis of current evidence and prices (Approve3 in Fig. 1). However, judging that research is worthwhile at this point is not sufficient for OIR guidance. An assessment of whether the benefits of early approval are expected to be greater than the opportunity costs of research that may be forgone to future patients is required. If the expected benefits are judged to be less than the opportunity costs, then OIR guidance would be appropriate (OIR1 in Fig. 1), whereas if they are judged to be greater, then approval would be appropriate (Approve2 in Fig. 1).

**Technologies not expected to be cost-effective**

A technology that is not expected to be cost-effective based on existing evidence is expected to impose negative NHE if it is approved. In these circumstances, Approve can be ruled out, but depending on the level of uncertainty in the current evidence and other changes that may occur, subsequent assessments and decisions are required before guidance is reached (see Fig. 1).

**Need for evidence.** Even if the technology is not expected to be cost-effective, the assessment may be uncertain; therefore, it remains a possibility that the technology might offer positive NHE. Again, the scale and consequences of uncertainty must be considered and whether additional research might potentially be worthwhile. If it is not, then the technology can be rejected (Reject4 in Fig. 1). Alternatively, if further research might be worthwhile, then an additional assessment is required of whether the type of evidence and research that is needed can be conducted without approval.

**Where research is possible without approval.** Generally, most types of research (including randomized clinical trials) will be possible without approval. The long-term value of research and the impact of other sources of uncertainty are required. If, following this reassessment, the expected benefits of research are judged to exceed the associated costs, then OIR would be appropriate (OIR2 in Fig. 1). Alternatively, if the costs of research are likely to exceed the longer term expected benefits, then the technology should be rejected at this point (Reject1 in Fig. 1).

**Where research is not possible without approval.** In some circumstances it is possible that certain types of evidence might be acquired only once a technology is in widespread use, for example, linking surrogates to longer term outcomes, longer term (or rare) adverse events, learning and incremental improvements, or identifying particular types of patients who might benefit most [27]. Where the type of research needed is not possible (or is
significantly more costly) without approval, the same assessment of the longer term benefits of research is required. If further research is judged not to be worthwhile following this reassessment, the technology can be rejected (Reject 2 in Fig. 1). Alternatively, if research is judged worthwhile, an additional assessment of whether the benefits of approval exceed the costs is required. In this case, approval will impose opportunity costs (negative expected NHE of widespread use of a cost-ineffective technology). The key question is whether the net benefits of the research exceed these opportunity costs. If they don’t, then the technology should be rejected even though research would have been worthwhile (Reject 3 in Fig. 1). Alternatively, if the net benefits of research more than offset the opportunity costs, then AWR would be appropriate even though the technology is expected to be cost-ineffective (AWR 2 in Fig. 1).

Therefore, AWR guidance for technologies not expected to be cost-effective is certainly possible but appropriate only in certain circumstances, where 1) the type of research needed is not possible without approval; 2) the long-term benefits of the research are likely to exceed the expected costs; and 3) the additional net benefits of such research exceed the opportunity costs of approving a cost-ineffective technology.

**Technologies with Significant Irrecoverable Costs**

Irrecoverable costs are those that once committed cannot be recovered if guidance is changed at a later date. Irrecoverable costs are most commonly thought of as “upfront” or capital costs of new facilities or equipment with long life expectancy. In most appraisals, these types of cost are first annuitized and then allocated pro rata to the number of patients likely to be treated during the lifetime of the equipment. If guidance remains unchanged throughout this period (i.e., research does not report or other sources of uncertainty resolve), then this common assumption has no influence. However, should guidance change (initial approval is withdrawn) before the end of the lifetime of the equipment, then, although future patents will no longer use the technology, the cost of the equipment that was allocated to them cannot be recovered.

Even in the absence of capital investment costs, most new technologies impose initial per patient treatment costs that exceed the immediate health benefits. These irrecoverable treatment costs are offset only by cost savings and health benefits in the longer run; that is, initially negative NHEs (losses) are only gradually compensated by later positive ones (gains). Therefore, a technology expected to be cost-effective may accumulate sufficient “gains” to compensate earlier “losses” only after some considerable time. Whether this type of irrecoverable opportunity cost is significant (i.e., might influence decisions) depends on whether treatment decisions for individual patients are irreversible, which, in part, depends on the nature of the disease. For example, in an acute condition, the decision to treat a particular patient with a technology cannot be reconsidered at a later date, whereas for a chronic condition the decision to treat can be delayed until the uncertainty is resolved. Therefore, the commitment of irrecoverable opportunity costs (negative NHE) can be avoided [21]. In these circumstances, OIR or Reject avoids this commitment and preserves the option to approve the technology at a later date when its purchase represents a “less risky investment” [21].
Technologies expected to be cost-effective
The presence of irrecoverable costs for a technology that is expected to be cost-effective will influence guidance and be regarded as “significant” only if there are future events (research reporting or other sources of uncertainty resolving) that might change guidance. For example, if research is possible with approval and is expected to be worthwhile, AWR does not necessarily follow as previously (e.g., see AWR in Fig. 1) because the impact of irrecoverable costs must also be considered. Now OIR may be more appropriate than AWR (e.g., the choice between OIR and AWR in Fig. 2), even though the research would be possible with approval because OIR avoids the commitment of irrecoverable costs until the results of research are known. This is especially the case when there are also other sources of uncertainty that might resolve while the research is being conducted because it increases the chance that guidance will be revised (e.g., OIR or AWR in Fig. 2).

If research is not possible with approval but is expected to be worthwhile, then OIR will be appropriate if the opportunity costs of early approval are judged to exceed the benefits (e.g., OIR rather than Approve in Fig. 2). Irrecoverable costs will tend to make OIR rather than approval more likely, particularly when there are other sources of uncertainty that might resolve while the research is being conducted (e.g., OIR rather than Approve in Fig. 2).

If research is not judged worthwhile, approval does not necessarily follow as previously (e.g., Approve in Fig. 1). Now the technology should be approved only if there are no other sources of uncertainty. If there are other sources of uncertainty, then an assessment of the benefits and costs of early approval that takes account of irrecoverable costs and the risk that guidance might change in the future is needed. Therefore,Reject rather than approval is possible, even though a technology is expected to be cost-effective, because the decision to commit the irrecoverable costs can be reconsidered once the other sources of uncertainty have resolved (e.g., Reject in Fig. 2).

Technologies not expected to be cost-effective
The presence of irrecoverable costs for technologies not expected to be cost-effective means that Reject is more likely to be appropriate than AWR when research is not possible without approval (see AWR in Fig. 3). This is because a decision to reject, although it may be revised to approve, generally does not commit irrecoverable costs. There may be circumstances when implementing guidance to reject a technology also requires resources if it has already been diffused into clinical practice. If these are significant they should be taken into account in the same way as other irrecoverable costs, tending to make AWR more likely to be appropriate.

Changes in the Effective Price of a Technology
Any change in the effective price of the technology will affect the key assessments, leading to different categories of guidance. For example, OIR for a technology, which is expected to be cost-effective, might be revised to Approve with a sufficient price reduction because the benefits of early approval will be greater and the value of additional evidence will tend to be lower (e.g., from OIR to Approve in Fig. 1). Similarly, AWR might be revised to Approve if the benefits of early approval now exceed the value of additional evidence (e.g., from AWR to Approve in Fig. 1). Therefore, consideration of the effect of price changes on OIR and AWR is needed when assessing the potential impact of discount schemes and more direct price negotiation.

Even if direct price negotiation is possible through a value-based pricing scheme[10], it will be important to retain OIR and AWR as possibilities because there is no guarantee that manufacturers will always agree to the lower price below which Approve rather than OIR or AWR would be appropriate, and there may be many circumstances when there is no effective price reduction that would make Approve appropriate. For example, Reject or OIR guidance may still be appropriate even if the effective price of a technology was zero if there is substantial uncertainty about its effectiveness and/or potential for harms.

Discussion
Each sequence of assessment and decision leads to a different category and type of guidance for technologies with differing characteristics, indications, and target populations. The different types of guidance illustrate how similar guidance might be arrived at but in different ways, adding to the transparency of the appraisal and helping to be explicit about the particular combination of considerations that underpin the guidance. There are many circumstances when AWR and OIR are applicable. For example, there are five types of OIR that may be appropriate when a technology is expected to be cost-effective. Even when research is possible with approval, OIR is appropriate if there are significant irrecoverable costs. Reject remains a possibility even for a cost-effective technology if there are irrecoverable costs. Therefore, the full range of categories of guidance (OIR and Reject as well as AWR and Approve) ought to be considered for technologies that on the balance of existing evidence and current prices are expected to be cost-effective. It is only approval that can be ruled out if a technology is not expected to be cost-effective; that is, cost-effectiveness is a necessary but not sufficient condition for approval but lack of cost-effectiveness is neither necessary nor sufficient for rejection. There are circumstances when AWR may be appropriate even when a technology is not expected to be cost-effective. Importantly, the category of guidance only partly depends on an assessment of expected cost-effectiveness. A number of other key assessments are required: the need for evidence; whether the type of research required is possible with approval; the expected longer term benefits and costs of research that is likely to be conducted; the impact of other sources of uncertainty that will resolve over time; and the significance of any irrecoverable costs.

This article has set out the key principles of what assessments are needed for coverage decisions. It has not specified how these assessments might be informed because this can differ across different jurisdictions. The explicit framework for OIR and AWR decisions is likely to have a number of implications for policy (e.g., drug pricing and incentives for evaluative research), as well as the process of appraisal (e.g., greater involvement of research commissioners) and the methods of appraisal (e.g., additional information, evidence, and analysis that might be required).

Coverage decisions that are based on some assessment of the likely health opportunity costs, whether or not a “threshold” is explicitly stated, sends a signal of how much health care systems can afford to pay for the benefits that new technologies offer. Manufacturers inevitably respond to these incentives through their pricing and investment decisions. Not only is price endogenous but the effects and the costs of developing and producing new technologies are endogenous as well. Therefore, the role of reimbursement agencies can be seen as signaling collective demand for the health benefits that new technologies offer, reflecting the social choices of how much resource is to be made available for health care. The endogeneity of price aligns the dynamic incentives for research and development with the needs and resource constraints of health care systems that reflect social values and mirror how temporary monopolists with patent protection respond to demand in other markets. The principles of how the need for evidence might influence coverage decisions and the implications that it has for pricing decisions by manufacturers enables greater subtlety in the signal sent and a better alignment of incentives for the development of new technologies and evaluative research available to support their use.
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