rare diseases, consideration of alternative acceptance criteria for public reimbursement /health service provision may be required.

The timing of cost-effectiveness over time and demonstrates the temporal impact on cost-effectiveness using a number of case-studies. The implications for decision making and market access are discussed. METHODS: A review of the factors that may influence cost-effectiveness over time and methodological approaches used to address these was conducted. Earlier analytical frameworks of studies from the 1990s in the fields of motor airbags, implantable cardiac defibrillators, statins, renal dialysis and hearing aids were revisited to re-estimate the cost-effectiveness. For example, parameters of an economic evaluation conducted in 1990 for erythropoietin were updated to 2004 values using a recent systematic review of clinical evidence together with revised unit costs and expert clinical opinion for resource utilisation. RESULTS: For the majority of case-studies examined, there was a trend for the reduction in cost-effectiveness ratios over time—e.g. for erythropoietin, the base-case cost per QALY decreased ten-fold over a 14 year period (£216,906 to £21,547). Significant factors included unit costs, dosage, utility gains and revised discounted rates. CONCLUSIONS: The timing of economic evaluation is critical in the estimation of cost-effectiveness. Production of this evidence may often be the first time that the conceptual framework of economic analysis has been applied to the technology, despite suggestions that economic evaluation should be used iteratively throughout the product life-cycle. This study has demonstrated that whilst there is a need for economic evaluation results to be timely to aid decision-making (i.e. at or around launch), it is important that the analysis is updated and reviewed periodically to assess whether cost-effectiveness has changed sufficiently to justify modifying the original decision.

EXAMINING THE QUALITY OF HEALTH ECONOMIC ANALYSES SUBMITTED TO THE REIMBURSEMENT AGENCIES IN SWEDEN AND FINLAND—A CROSS COUNTRY COMPARISON

Engstrom A, Kivioja A

1Pharmaceutical Benefits Board, Solna, Sweden; 2Ministry of Social Affairs and Health, Helsinki, Finland

OBJECTIVES: To compare the quality of the health economic material submitted to the Swedish Pharmaceutical Benefits Board and the Finnish Pharmaceuticals Pricing Board as part of the application for reimbursement for new pharmaceuticals. METHODS: The health economic evaluations were reviewed in each country against two checklists, marking each question Yes/No/Not Applicable. The checklists used were: 1) the respective national Guidelines transformed into yes or no questions, and 2) the QHES check list, a validated instrument, was also used to provide a common comparator. The central estimate of cost effectiveness was collected (cost per QALY) as well as whether the application was accepted or rejected in each country. RESULTS: The Swedish scores range from 0.24 to 0.87 and on the QHES from 0.09 to 1, with a mean quality of 0.61 and 0.67 respectively. The Finnish scores range from 0.58 to 0.96 and on the QHES from 0.28 to 0.84, with a mean quality of 0.76 and 0.62 respectively. The correlation between the respective national guidelines and the QHES scores is modest (approx. 0.7 both in Sweden and in Finland). This is mostly due to country specific criteria. There was a low observed correlation between quality score and acceptance in Sweden and also in Finland. Likewise, the correlation between cost per QALY and decision to accept/reject is low to medium. CONCLUSIONS: Health economic material as part of applications to reimbursement agencies varies widely in quality. There are differences even for the same product in the two countries. Secondly, due to the relatively small number of applications studied and the even fewer rejections, it is difficult to draw firm conclusions regarding the value the pricing authorities studied place on a QALY.

PAYMENT FOR PHARMACY HANDLING COSTS IN THE HOPD: 2006 REIMBURSEMENT IMPLICATIONS

Baker II, Panirites C

The Resource Group, Dallas, TX, USA; 2Scios, Inc, Fremont, CA, USA

OBJECTIVE: Medicare reform legislation requires the Medicare Payment Advisory Commission (MedPAC) to conduct a study of hospital pharmacy handling costs and to recommend whether payment for handling costs of Part B specified outpatient drugs should be made under the Medicare Hospital Outpatient Prospective Payment System (OPPS). This study explores the MedPAC approach to pharmacy departmental costs and potential reimbursement implications. METHODS: The MedPAC rationale and recommendations were deconstructed, evaluated, and compared to data about pharmacy operations obtained from two studies. The first study collected data from on-site observations of infusion therapy and related pharmacy activities in 24 hospital outpatient departments (HOPDs) located in 19 states. A subsequent telephone survey gathered information about staffing and workflow from 30 other hospital pharmacy directors located in 16 states and the District of Columbia. RESULTS: The MedPAC report places pharmacy costs into five categories; concludes that hospitals can estimate these costs; and recommends that hospitals develop and submit charges for a new set of handling fee APCs. Findings from the comparative on-site observations and telephone survey, however, reveal that significant portions of certain costs are not charged to the pharmacy. For example, 69.1% of respondents had information systems costs charged outside the department while 34.5% reported their entire information systems costs charged elsewhere. Likewise, 73.3% of respondents reported reimbursement, chargemaster, clerical and transport staff utilized by the pharmacy department but not charged to the department. Such disparities in departmental costing will significantly hinder the uniform interpretation of costs assumed by MedPAC. CONCLUSIONS: Payment methods in 2006 for handling costs of Part B separately paid drugs will be derived in large part from the MedPAC recommendations. If the payment methodology does not take existing variations in recording pharmacy expense outside the department, hospital providers may well be underpaid for handling costs in 2006.