Long-term effectiveness and cost-effectiveness of screening for Hepatitis C virus infection

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Background: Hepatitis C virus (HCV) infection is an emerging problem in public health. In most countries, the majority of HCV infected people are yet undiagnosed. Early detection and treatment may result in better health outcomes and save costs by preventing future advanced liver disease. The evidence for long-term effectiveness and cost-effectiveness of HCV screening was systematically reviewed. Methods: We performed a systematic literature search on long-term health-economic effects of HCV screening and included Health Technology Assessment (HTA) reports, systematic reviews, long-term clinical trials, full health economic and decision-analytic modelling studies with a sufficiently long time horizon and patient-relevant long-term outcomes such as life-years gained (LYG) or quality-adjusted life years (QALY) gained. Economic results were converted to 2005 Euros. Results: Seven studies were included. Target population, HCV prevalence, study perspective, discount rate, screening and antiviral treatment mode varied. The incremental effectiveness of HCV screening and early treatment compared to no screening and standard care varied from 0.0004 to 0.066 LYG, and from 0.0001 to 0.072 QALY. Incremental cost-effectiveness and cost-utility ratios of HCV screening vs. no screening were 3900–243 700 €/LYG and 18 300–151 000 €/QALY. HCV screening seems to be cost-effective in populations with high HCV prevalence, but not in low HCV prevalence populations. Conclusions: HCV screening and early treatment have the potential to improve average life-expectancy, but should focus on populations with elevated HCV prevalence to be cost-effective. Further research on the long-term health-economic impact of HCV screening when combined with appropriate monitoring strategies in different European health care systems is needed.

Keywords: chronic hepatitis C, cost effectiveness, screening.

Introduction

Chronic Hepatitis C (CHC) is an emerging problem in public health. In Europe, the Hepatitis C virus (HCV) infection affects > 1% of the population with a HCV-incidence of 8.6/100 000.¹,² HCV prevalence differs considerably across countries and risk groups.³ The highest HCV prevalence (36–81%) is currently found in intravenous drug users (IDUs).¹

The majority of HCV-infected people progress to chronic disease.⁴ Approximately 15–20% of CHC cases develop cirrhosis within 20–30 years,⁵–¹² which is associated with a high risk for advanced liver disease, quality of life impairment, reduced life expectancy and high treatment costs. CHC is considered to be the leading cause of liver cancer and liver transplantation in Europe.¹³

Screening for CHC clearly fulfils the general criteria for population screening¹⁴,¹⁵ and may help to identify HCV-infected patients in an early stage of the disease (e.g. mild chronic hepatitis without fibrosis), so that they can be adequately monitored and treated. Moreover, it has been reported that it may be cost-effective to treat patients diagnosed with mild disease.¹⁶,¹⁷ Furthermore, for the majority of acute HCV cases, which present no symptoms, early treatment and for symptomatic acute HCV cases watchful waiting may be currently the most effective and cost-effective strategies.¹⁸ Thus, early detection and early treatment may have the potential to result in better health outcomes and to save costs by preventing future advanced liver disease. Another important reason to identify unaware HCV-infected persons is to prevent further HCV-transmission using appropriate interventions to change behaviour leading to HCV transmission (e.g. needle sharing).

However, currently most European countries lack specific policies for HCV screening. Only few European countries perform HCV screening in special subpopulations with elevated HCV prevalence. But even in these cases, the recommendations and medical practices are heterogeneous.¹⁹–²¹ In March 2007, the European Parliament called for EU-wide action on Hepatitis C by formally adopting the Written Declaration on Hepatitis C.²² Specifically, the European Parliament calls for a council recommendation on Hepatitis C screening to ensure early diagnosis and wider access to treatment and care within the member states. Furthermore, the European Liver Patients Association (ELPA) strongly suggests that the European Union should encourage tailored screening campaigns that target people in at-risk groups.²³

Despite all potential benefits, HCV screening may have substantial health-economic consequences and it is not clear whether it leads to improved long-term health outcomes, because not all CHC patients will develop progressive liver disease in their lifetime, and not all CHC patients benefit from antiviral treatment.¹⁶,²⁴,²⁵ Furthermore, current antiviral...
treatment options are costly, and impose the burden of side effects.\textsuperscript{16,24,25} Therefore, a thorough assessment of HCV screening must consider all consequences for individuals and society during a sufficiently long time horizon.

In this review, we systematically evaluated the current evidence on long-term effectiveness and cost-effectiveness of screening for Hepatitis C virus infection in different populations.

Methods

A systematic literature search was conducted using the databases Medline, Cochrane Database of Systematic Reviews, Cochrane central register of controlled trials (CENTRAL), and the NHS databases abstracts of reviews of effects (DARE), Health technology assessment (HTA) and Economic evaluation database (NHS EED) to identify studies assessing the clinical and economic long-term consequences of screening for Hepatitis C virus infection (HCV). The time horizon of the literature search was limited to March 2007. All references were imported into a literature database using a literature management software program (EndNote 9.0, Thomson ResearchSoft TM, Thomson Corporation, Stamford, CT, USA).

First, reference titles and abstracts were screened for relevant articles. In a second step, studies were selected based on a priori inclusion and exclusion criteria after reading the full text document. We included health technology assessment (HTA) reports, systematic reviews, long-term clinical trials, full health economic studies and decision-analytic modelling studies assessing the impact of screening for Hepatitis C virus infections. As clinical and economic consequences of screening occur over a long time horizon, we only included studies that reported both long-term effectiveness and cost-effectiveness in terms of life-years gained (LYG), quality-adjusted life-years gained (QALY), lifetime cost per life-year gained (Cost/LYG) or cost per quality-adjusted life-year gained (Cost/QALY).

We excluded studies in languages other than English or German, editorials, letters, abstracts, unsystematic reviews, studies reporting only short-term effectiveness data (e.g. sustained virological response, SVR), studies assessing screening of blood donations or serological testing during antiviral treatment. We also excluded studies that did not report sufficient data to derive incremental effectiveness and cost-effectiveness ratios or cost-effectiveness studies reporting only costs per HCV case detected.

We systematically extracted the results from the publications and summarized the information in evidence tables reporting clinical and economic outcomes.

If necessary and possible, we recalculated the incremental cost-effectiveness ratios (ICER) or incremental cost-utility ratios (ICUR) from the data reported in the publication. To facilitate comparison across countries and to enable other countries to transfer our results into their currencies, all costs were converted to 2005 Euro (€) using gross domestic product purchasing power parities (GDP PPP\textsuperscript{26}) (conversion to Euro of the index year) and the German Consumer Price Index (CPI) (inflation to the year 2005).\textsuperscript{26,27} Germany was used as the reference country for the cost conversion, because it is the country with the largest population in Europe.\textsuperscript{28}

Results

Literature search

A total of 127 unique references were retrieved. Ten publications,\textsuperscript{20,29–37} including two HTA reports\textsuperscript{20,36} assessing lifetime health effects and costs of screening for Hepatitis C met the inclusion criteria. No long-term clinical trial assessing the long-term effectiveness (e.g. mortality) of screening for Hepatitis C virus infection and early HCV-treatment was identified.

Two publications by Stein\textsuperscript{33,34} et al. reported the cost-effectiveness results of a decision-analytic model performed within an HTA report conducted by the National Institute for Health and Clinical Excellence (NICE).\textsuperscript{20} Thompson Coon\textsuperscript{37} et al. reported the cost-effectiveness results of a decision-analytic model performed within an HTA report conducted by the NHS R&D HTA Program.\textsuperscript{36} Only the original data from the HTA reports were considered, leaving seven studies in the review.

Long-term effectiveness

In the absence of clinical trials, meta-analyses and health technology assessment reports evaluating the long-term effectiveness of HCV screening, we based our results on decision-analytic modelling studies that included an analysis of long-term effectiveness of screening for Hepatitis C virus infection and early HCV-treatment in terms of undiscounted life years and/or quality-adjusted life years gained compared to no screening and standard care.

Five out of seven cost-effectiveness studies reported undiscounted life years and/or quality-adjusted life years gained for screening and early HCV-treatment compared to no screening and standard care (table 1).\textsuperscript{20,29,30,35,36}

The values for life years gained due to screening and early treatment varied from 0.0004 LYG (0.15 life days) for screening blood recipients to 0.066 LYG (24.1 life days) for screening all patients assessed for HBV vaccination attending drug and alcohol services. QALYs varied from no gain for screening in pregnant women to 0.072 QALYs (i.e. 26 quality-adjusted life days) for screening in patients assessed for HBV vaccination attending drug and alcohol services. Screening in populations with elevated HCV prevalence (e.g. IDU) was more effective in terms of life-years or QALYs gained. Studies reported 0.036–0.066 LYG (13.1–24.1 life days) for populations with 42–68% HCV prevalence (0.010–0.072 QALYs/3.7–26.3 quality-adjusted life days; 32–68% HCV prevalence) vs. 0.0004–0.013 LYG (0.1–4.7 life days) for populations with 3–16% HCV prevalence (0–0.022 QALYs/0–8.0 quality-adjusted life days; 1–16% HCV prevalence).

Long-term cost-effectiveness

Health technology assessment reports

Two HTA reports were included. One summarized results from economic studies evaluating HCV-screening programmes, and both HTA reports conducted a cost-effectiveness analysis.

Stein\textsuperscript{20} et al.\textsuperscript{20} systematically reviewed the evidence from health economic studies evaluating HCV-screening programmes. All reviewed studies had methodological limitations and the results were of limited transferability to the UK context. Based on their decision-analytic results, the authors concluded that screening for Hepatitis C in intravenous drug users in contact with medical services may be moderately cost-effective. However, the authors recommend interpreting their results with caution because of substantial uncertainty around the acceptability of screening, the adherence to treatment and the simple nature of the model. General screening in genito-urinary medicine (GUM) clinics is less cost-effective and associated with greater uncertainty than screening IDUs in contact with medical services.
<table>
<thead>
<tr>
<th>Study, Country</th>
<th>Population</th>
<th>HCV prevalence (%)</th>
<th>Screening/Treatment</th>
<th>Incremental life years (LYG)</th>
<th>Incremental quality-adjusted life years (QALY)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Castelnuovo et al.(^{16}) (Thompson Coon et al.(^{17})) NHS R&amp;D HTA Programme UK</td>
<td>Former IDUs, mean age 37 years</td>
<td>49</td>
<td>Systematic screening vs. no systematic screening (spontaneous presentation to screening possible); HCV-positives receive treatment: PegIFN+RBV</td>
<td>0.058</td>
<td>0.071</td>
</tr>
<tr>
<td></td>
<td>General practice, mean age 37 years</td>
<td>12.5</td>
<td></td>
<td>0.010</td>
<td>0.017</td>
</tr>
<tr>
<td></td>
<td>Former and current IDUs in general practice, mean age 37 years</td>
<td>49</td>
<td></td>
<td>0.036</td>
<td>0.071</td>
</tr>
<tr>
<td></td>
<td>All patients assessed for HBV vaccination attending drug and alcohol services, mean age 37 years</td>
<td>68</td>
<td></td>
<td>0.066</td>
<td>0.072</td>
</tr>
<tr>
<td></td>
<td>Prisoners at reception, mean age 37 years (general counseling)</td>
<td>16</td>
<td></td>
<td>0.013</td>
<td>0.022</td>
</tr>
<tr>
<td></td>
<td>Prisoners at reception, mean age 37 years (counseling with IDU focus)</td>
<td>42</td>
<td></td>
<td>0.036</td>
<td>0.058</td>
</tr>
<tr>
<td>Jusot and Colin(^{18}) France</td>
<td>Blood recipients &lt; 40 years</td>
<td>3</td>
<td>Screening with EIA3 after transfusion, treatment for HCV-positives with Knodell score 5: IFN vs. no screening + no medical therapy</td>
<td>0.0085</td>
<td>n.a.</td>
</tr>
<tr>
<td></td>
<td>Blood recipients 40–65 years or receiving low-volume transfusions or hospitalized in a surgery department</td>
<td>3</td>
<td>Screening with EIA3 after transfusion, treatment for HCV-positives with Knodell score 5: IFN vs. no screening + no medical therapy</td>
<td>0.0004</td>
<td>n.a.</td>
</tr>
<tr>
<td></td>
<td>Blood recipients receiving high-volume transfusions</td>
<td>3</td>
<td>Screening with EIA3 before and after transfusion, (treatment same as above)</td>
<td>0.0030–0.0047(^{a})</td>
<td>n.a.</td>
</tr>
<tr>
<td>Leal et al.(^{29}) UK</td>
<td>IDUs in contact with drug services, mean age: n.a.</td>
<td>60</td>
<td>Screening vs. no screening: HCV-positives with moderate to severe CHC receive treatment: IFN</td>
<td>n.a.</td>
<td>0.015(^{b})</td>
</tr>
<tr>
<td>Plunkett et al.(^{35}) USA</td>
<td>Pregnant women, mean age 30 years</td>
<td>1</td>
<td>Screening vs. no screening: 70% (screened) or 20% (unscreened) of HCV-positives with moderate CHC receive treatment: PegIFN + RBV</td>
<td>n.a.</td>
<td>–0.00011</td>
</tr>
<tr>
<td>Stein et al.(^{20}) (Stein et al. 2003(^{33,34})) NHS R&amp;D HTA Programme UK</td>
<td>IDUs in contact with drug services, mean age 32 years</td>
<td>32</td>
<td>Screening and treatment as above plus Caesarian delivery</td>
<td>n.a.</td>
<td>0.0001</td>
</tr>
<tr>
<td></td>
<td>Genito-urinary medicine clinic attendees, mean age 36 years</td>
<td>1.5</td>
<td>Screening vs. no screening: 50% of HCV-positives with moderate CHC receive treatment: IFN + RBV</td>
<td>n.a.</td>
<td>0.01003(^{2})</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Screening vs. no screening: 50% of HCV-positives with moderate CHC receive treatment: IFN + RBV</td>
<td>n.a.</td>
<td>0.0004(^{2})</td>
</tr>
</tbody>
</table>

\(^{a}\): Range reported in the original study for the first, second and third year

\(^{b}\): Calculated with data in the original publication

n.a. = not available; IFN = interferon; RBV = ribavirin; EIA3 = enzyme linked assay third generation; PCR = polymerase chain reaction.
Castelnuovo et al. performed a decision-analytic cost-effectiveness study to evaluate screening (named ‘case-finding’) in patients attending general medical practice or special drug and alcohol services, and in prisoners at reception with a focus on former IDUs. Based on their analyses, the authors concluded that screening in these target populations is likely to be cost-effective despite some uncertainty around the acceptance of testing and treatment.

Cost-effectiveness studies

Seven cost-effectiveness studies evaluating HCV screening in different population settings were included in our review (table 2). Three studies were conducted in the UK, two in France, and two in the USA. Studies varied in terms of target population, study perspective, time horizon, discount rate and compared strategies including screening and antiviral treatment mode.

Five studies evaluated average costs at risk for Hepatitis C (HCV prevalence 1–3.8%). Of those one study evaluated HCV screening in asymptomatic, average-risk adults in the USA, one study examined screening in the general French population, and another study assessed screening in pregnant women in the USA. Two studies considered screening in blood recipients and one in general Genito-urinary medicine clinic attendees.

Four studies assessed the cost-effectiveness of screening in different populations at higher risk for HCV (HCV prevalence 7–80%). Four studies evaluated HCV screening in populations with a history of IDU in different settings, two studies assessed the cost-effectiveness of general screening in attendees of special medical services, and one evaluated HCV screening in prisoners at reception.

Most studies compared systematic screening (and antiviral treatment for detected HCV-positives) to no-systematic screening, allowing for the possibility of spontaneous case detection with subsequent antiviral treatment. The percentage of HCV positives eligible for treatment varied. Some studies compared screening and antiviral treatment for detected HCV-positives to no screening and no treatment. The antiviral treatment regimens (interferon/interferon plus ribavirin/peginterferon plus ribavirin) and algorithms (e.g. treat all HCV-positives or only those with severe liver histology) varied. Only three studies evaluated screening followed by peginterferon plus ribavirin, the current recommended standard antiviral therapy.

The incremental cost-effectiveness ratios (ICER) of HCV screening vs. no screening varied across countries about the cost-effectiveness threshold. To give an idea of the numbers reflect the average incremental life expectancy per person screened. This translates to many persons with no gain and some persons with several years or decades gain in life expectancy.

Cost-effectiveness ratios compared to more targeted screening (e.g. screening only IDUs in these settings).

Figure 1 shows the incremental ICER and ICUR ratios of screening for different HCV prevalence and different antiviral treatment strategies. Most studies evaluated the ICERS/ICURs in populations with HCV prevalence above 10%. Only four studies reported results for populations with a lower HCV prevalence. Many studies evaluated screening followed by antiviral treatment with interferon or interferon plus ribavirin, which are not current standard treatment options anymore. Peginterferon plus ribavirin, the recommended standard antiviral treatment yields more LYs/QALYs gained and results in much lower ICERS/ICURs. Therefore, figure 1c and d shows ICERS/ICURs for screening followed by treatment with peginterferon plus ribavirin, only. The majority of these studies reported ICURs below 40 000 €/QALY gained (ICER: 50 000 €/LYG) in populations with HCV prevalence above 10%, and higher ICURs (77 000–1 150 000 €/QALY gained) in low HCV prevalence populations (results from two studies).

Discussion

We performed a systematic review on the long-term effectiveness and cost-effectiveness of screening for HCV infection.

Depending on HCV prevalence and risk selection mode, the incremental long-term effectiveness of HCV screening and early treatment compared to no screening and standard care varied from 0.0004 LYG (0.15 life-days gained) to 0.066 LYG (24 life-days gained) and from 0.0001 QALY (0.04 quality-adjusted life-days gained) to 0.072 QALY (26 quality-adjusted life-days gained). To put these figures into perspective, they can be compared with other screening programs. For example, biennial cervical cancer screening compared to no screening is associated with a gain of 92 life days. Moving from a 2-year to a 1-year interval is associated with a gain of four life days.

Given 1% undetected HIV-prevalence, one-time HIV screening in US health care settings was reported to increase life-expectancy by 3.9 days (2.9 quality-adjusted life days). Screening every 5 years would gain additional 0.97 days (0.70 quality-adjusted life days). It must be noted, that these numbers reflect the average life expectancy per person screened. This translates to many persons with no gain and some persons with several years or decades gain in life expectancy.

The incremental cost-effectiveness ratios varied over a wide range depending on target population (e.g. HCV prevalence, age, etc.), study perspective, time horizon, discount rate and compared strategies including screening settings and antiviral treatment strategies. Therefore, the comparability of the results is limited.

HCV screening vs. no screening resulted in ICURs ranging from 18 300 to 1 151 000 €/QALY, if screening was not dominated. In the reviewed studies, HCV screening was considered cost-effective (ICURs below 40 000 €/QALY for treatment with peginterferon plus ribavirin) in populations with an elevated HCV prevalence such as intravenous drug users. General HCV screening in average-risk adults was unlikely to be effective and cost-effective.

However, cost-effectiveness should not be the main criterion for the decision to implement HCV screening. Given the substantial number of prevalent iatrogenic HCV-infected cases other ethical concepts such as fairness and equity may be considered as well.

Cost-effectiveness is depending on the willingness-to-pay in a certain society, which depends on several economical, social and political factors. There is currently no general agreement across countries about the cost-effectiveness threshold. To give a measurement on the incremental cost-effectiveness ratios of
<table>
<thead>
<tr>
<th>Study, Country</th>
<th>Currency, Year, Perspective</th>
<th>Discount rate (%)</th>
<th>Comparator</th>
<th>Target Population</th>
<th>ICER (€/LYG)</th>
<th>ICUR (€/QALY)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Castelnuovo et al.16 (Thompson Coon et al.17)</td>
<td>UK£, 2004, National Health Services (NHS)</td>
<td>6 (costs), 1.5 (effects)</td>
<td>Systematic screening vs. no systematic screening (spontaneous presentation to screening possible); HCV-positives receive treatment: PegIFN + RBV</td>
<td>Former IDUs, general case, mean age 37 years, 49% HCV prevalence</td>
<td>30232</td>
<td>24 858</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>General practice, mean age 37 years, 12.5% HCV prevalence</td>
<td>38633</td>
<td>23 321</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Former and current IDUs in general practice, mean age 37 years, 49% HCV prevalence</td>
<td>30194</td>
<td>24 827</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td>All patients assessed for HBV vaccination attending drug and alcohol services, mean age 37 years, 68% HCV prevalence</td>
<td>28689</td>
<td>26 365</td>
</tr>
<tr>
<td>NHS R&amp;D HTA Programme</td>
<td>UK</td>
<td></td>
<td>(Screening included general lecture on HCV) (Screening included lecture with focus on IDU and risk of HCV)</td>
<td>Prisoners at reception, mean age 37 years, 16% HCV prevalence</td>
<td>50833</td>
<td>30 231</td>
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<td>Prisoners at reception, mean age 37 years, 42% HCV prevalence</td>
<td>40301</td>
<td>24 813</td>
</tr>
<tr>
<td>Jusot and Colin20</td>
<td>FF, 1996, Health care system, 30 years time horizon</td>
<td>No discount rate</td>
<td>Screening with EIA3 after transfusion, treatment for HCV-positives with Knodell score 5: IFN vs. no screening + no medical therapy</td>
<td>Blood recipients &lt;40 years, 3% HCV prevalence</td>
<td>140674</td>
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<td></td>
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<td></td>
<td>Screening with EIA3 after transfusion, treatment for HCV-positives with Knodell score 5: IFN vs. no screening + no medical therapy</td>
<td>Blood recipients 40-65 years or receiving low-volume transfusions or hospitalized in a surgery department, 3% HCV prevalence</td>
<td>477654</td>
<td>—</td>
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<tr>
<td></td>
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<td></td>
<td>Screening with EIA3 before and after transfusion, (treatment same as above)</td>
<td>Blood recipients receiving high-volume transfusions, 3% HCV prevalence</td>
<td>144970</td>
<td>—</td>
</tr>
<tr>
<td>Leal et al.29</td>
<td>UK, 1997, n.a.</td>
<td>6</td>
<td>Screening vs. no screening; HCV-positives with moderate to severe CHC receive treatment: IFN (IFN + RBV)</td>
<td>IDUs in contact with drug services, 60% HCV prevalence</td>
<td>—</td>
<td>119 754 (18 267-34537*)</td>
</tr>
<tr>
<td>Loubiere et al.32</td>
<td>FF, 1998, Health care system</td>
<td>3</td>
<td>Screening with EIA3+EIA3 vs. no screening + no treatment; 50% of CHC cases and 40% of cirrhosis cases receive treatment: IFN + RBV</td>
<td>IDUs, 80% HCV prevalence</td>
<td>3881</td>
<td>—</td>
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<td></td>
<td></td>
<td></td>
<td>Screening with EIA3+PCR vs. no screening + no treatment; treatment as above</td>
<td>—</td>
<td>9742</td>
<td>—</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Screening with EIA3+PCR vs. no screening + no treatment; treatment as above</td>
<td>Patients transfused before 1991, 7% HCV prevalence</td>
<td>—</td>
<td>Dominated by EIA3+PCR 243737</td>
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<td></td>
<td></td>
<td></td>
<td>Screening with EIA3+PCR vs. no screening + no treatment; treatment as above</td>
<td>General French population, 1.2% HCV prevalence</td>
<td>—</td>
<td>Dominated by EIA3+PCR 5005</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Screening with EIA3+PCR vs. no screening + no treatment; cirrhosis treatment as above</td>
<td>—</td>
<td>—</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Screening with EIA3+PCR vs. no screening + no treatment; cirrhosis treatment as above</td>
<td>—</td>
<td>—</td>
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<tr>
<td>Plunkett et al.35</td>
<td>US$, 2003, Health care system</td>
<td>3</td>
<td>Screening vs. no screening; HCV-positives receive treatment: PegIFN + RBV</td>
<td>Pregnant women, 1% HCV prevalence</td>
<td>—</td>
<td>No screening dominant</td>
</tr>
<tr>
<td>Singer et al.31</td>
<td>US$, 2001, Societal</td>
<td>3</td>
<td>Screening vs. no screening: 20% of HCV-positives receive treatment: IFN + RBV</td>
<td>Asymptomatic, average risk adults, mean age 35 years, 3.8% HCV prevalence</td>
<td>—</td>
<td>1 150 976</td>
</tr>
<tr>
<td>Stein et al.30 (Stein et al.33,34)</td>
<td>UK£, 2001, National Health Services (NHS)</td>
<td>6 (costs), 1.5 (effects)</td>
<td>Screening vs. no screening: 50% of HCV-positives with moderate CHC receive treatment: IFN + RBV (PegIFN + RBV)</td>
<td>IDUs in contact with drug services, mean age 32 years, 32% HCV prevalence</td>
<td>—</td>
<td>46 707 (23598)</td>
</tr>
<tr>
<td>NHS R&amp;D HTA Programme</td>
<td>UK</td>
<td></td>
<td>All screened, 50% of HCV-positives with moderate CHC receive treatment: IFN + RBV (PegIFN + RBV)</td>
<td>Genito-urinary medicine clinic attendees, mean age 36 years, 1.5% HCV prevalence</td>
<td>—</td>
<td>140 471 (77 052)</td>
</tr>
<tr>
<td></td>
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<td></td>
<td>IDUs screened, 50% of HCV-positives with moderate CHC receive treatment: IFN + RBV</td>
<td>—</td>
<td>45 076</td>
<td></td>
</tr>
</tbody>
</table>

a: Depending on treatment duration (24 or 48 weeks) and ribavirin dosage (1000 or 1200mg/d)
n.a. = not available; IFN = interferon; RBV = ribavirin; EIA3 = enzyme linked assay third generation; PCR = polymerase chain reaction.
well-accepted screening programs, cytological screening for cervical cancer every 3 years compared to no screening costs £1800 per life year gained in the UK, £14 000 €/LYG in Germany, and 8400 US$/LYG in the USA. Screening blood donors for HIV costs 14 000 US$/LYG. Given 1% undetected HIV-prevalence, one-time HIV screening in US health care settings would result in 41 700 US$/QALY, screening every 5 years 123 600 US$/QALY. In the absence of long-term clinical trials, all results were retrieved from decision-analytic studies, which link diagnostic and clinical short-term outcomes (e.g. test sensitivity and specificity or viral response) to clinical long-term outcomes (e.g. mortality and long-term quality of life). The included studies were heterogeneous in regard to health economic analysis techniques (e.g. time horizons, discounting, etc.), HCV population prevalence, acquisition risk factors and antiviral therapy. Therefore, the outcomes in terms of life years gained, quality adjusted life years and incremental cost effectiveness ratios varied over a wide range. However, several results were logical and predictable, for example, screening is more cost effective in higher prevalence or higher risk populations—a result that has been reported for other diseases, too.

Like all decision-analytic models, screening models must simplify the real world for more transparency and the possibility to analyse specific research questions. However, some methodological and structural model assumptions may have an important impact on clinical and economic outcomes and could lead to bias in favour for or against HCV screening. Thus, it is important to discuss some aspects essential for a valuable screening model.

First, it is important to allow for the possibility of spontaneous case detection by symptoms with subsequent antiviral treatment in the non-screening strategy of any HCV-screening model. Without these estimates, the benefits of the screening strategy are overestimated and outcomes are biased in favour of the HCV-screening strategy. Second, the setting of antiviral treatment in both strategies is very important. No treatment in the non-screening strategy or ‘wait and treat cirrhosis’ vs. ‘screen and treat all HCV-positive patients’ may overestimate both the incremental benefits and costs of screening. Therefore, antiviral treatment should be considered for chronic HCV-patients (detected through screening, symptoms or spontaneous presentation) in both strategies according to recent treatment guidelines. Third, most studies considered antiviral treatment with interferon plus ribavirin, and two studies used even interferon monotherapy. Only three studies considered peginterferon plus ribavirin. Having better treatment options and administering antiviral treatment according to genotype-specific guidelines with early treatment stop for patients not responding would allow tailoring treatment efficiently, which
would reduce adverse effects, harms, and antiviral treatment costs, and improve the cost-effectiveness of HCV screening due to better clinical and economic outcomes. Fourth, eligibility of patients for and adherence to antiviral treatment should be considered. In particular, any HCV-screening model should consider a ‘wait and see’ strategy in the screening arm, because not all patients necessarily should or want to be treated immediately after HCV detection.50 HCV screening and watchful monitoring HCV-infected patients may be more effective and cost-effective than screening with immediate treatment of all HCV-infected patients, since a fraction of HCV-infected patients may not develop fibrosis or cirrhosis during their lifetime. This is particularly important for the elderly.

The age at which HCV-infected patients are identified and treated is a very important modelling factor as well. Most studies used an average age of 40 years for the evaluated population, which may be adequate for patients with CHC, which already developed symptoms. However, HCV screening may detect HCV-infected individuals earlier at significantly younger age. In addition certain HCV-infected populations at risk for advanced liver disease, such as intravenous drug user or ethnic minority groups, who have acquired HCV iatrogenically in early childhood, have a significantly younger average age. Thus, in these cases benefits from early detection and treatment may be underestimated.

Discounting costs and effects is important and affects ICERs, since the clinical and economic benefits of screening due to avoided cirrhosis and its sequelae occur in the distant future, whereas the costs of screening and antiviral treatment occur much earlier. One study30 in France did not discount at all and two studies 20,36 conducted in the UK used different discount rates for costs and effects. The ICERs increased significantly in sensitivity analyses when cost and effects were equally discounted with 3.5% annually (e.g. from 16 514 £/QALY to 33 235 £/QALY56).

Most studies used a lifelong time horizon for their analyses which is the most adequate timeframe to use. As benefits that occur far in the future will not be considered within shorter time horizons, estimated cost-effectiveness ratios may be too high. One study used a 30 year time horizon.30 As cirrhosis and its complications develop slowly within 10–30 years, even this time horizon may be too short and benefits may be underestimated.

All studies included in this review take into account the natural history of chronic Hepatitis C disease progression and mortality from CHC-related complications. Only one study used the natural history of chronic Hepatitis B disease progression, as at that time no information existed regarding Hepatitis C progression. However, it was not always clear whether slower progression rates were considered for screened populations tending to present histological milder Hepatitis C compared to non-screened populations mostly detected by symptoms. Several studies reported that patients with mild CHC and normal ALT levels may have a reduced risk of progression to cirrhosis compared to patients with more severe histology or elevated ALT levels,7,51–54 Furthermore, analyses for CHC patients co-infected with HIV should assume higher progression rates to CHC-related liver diseases than analyses in non-co-infected CHC patients,55,56.

In addition, most studies used age- and gender-specific mortality rates of the general population for the background mortality for CHC patients. However, background mortality is often higher due to co-morbidity from other diseases such as HIV- or HBV-coinfection, or in case of IDUs from continuation of or relapse to drug abuse. Even patients with moderate CHC or cirrhosis that respond to antiviral treatment continue to have an increased risk of developing hepatocellular carcinoma, which is associated with significant mortality.

Overall, this review discovered many study limitations and the need for further systematic research in HCV screening. Particularly, health-economic studies in population with low or average HCV prevalence evaluating HCV screening combined with different strategies of monitoring and antiviral treatment of HCV-positives according to current treatment standard are required.

Finally, it must be mentioned, that due to different epidemiology, health care systems, disease management practice patterns and treatment costs in different European countries, results cannot be generalized and are difficult if not impossible to be directly transferred from one country to another. Further research should focus on the development of a Pan-European Hepatitis C screening model that fulfils the quality criteria discussed above and which can be adapted to the context of the different health care systems and countries within Europe.

**Conclusion**

Although HCV screening fulfils general population screening criteria, specific well-formulated national programs for Hepatitis C screening are lacking in most European countries. Based on current evidence, HCV screening and early treatment has the potential to improve average life-expectancy, but should focus on populations with elevated HCV prevalence to be cost-effective. Further research is needed to investigate the long-term health-economic impact of HCV screening when combined with appropriate monitoring and treatment strategies in different European health care systems. Further assessments should focus on determining optimal target groups and settings that yield effective and cost-effective HCV screening strategies.

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Key points

- Although HCV screening fulfils general population screening criteria, specific well-formulated national public health programs for hepatitis C screening are lacking in most European countries.
- According to this review, HCV screening with early treatment has the potential to improve average life-expectancy, but should focus on populations with elevated HCV prevalence to be cost-effective. Appropriate target groups could be selected based on risk factor profiles.
- Appropriate monitoring and treatment strategies for detected early disease may improve the cost-effectiveness of HCV screening.
- In view of the multitude of iatrogenic infections, however, cost-effectiveness may not be the only decision criterion for the implementation of HCV screening. Aspects like fairness might be considered as well.
- Further research should focus on the public-health impact of HCV screening when combined with appropriate monitoring and treatment strategies and on determining optimal target groups and settings.

References


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